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(54) Title: BRASSICA POLYMORPHISMS

(57) Abstract

The invention provides oligonucleotides and their complements that can be used as allele-specific probes or primers for sequencing, oligonucleotide probe hybridization, and allele-specific amplification. Such oligonucleotides can be used, for example, to facilitate genetic distinction between individual plants in plant populations.

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BRASSICA POLYMORPHISMS

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CROSS-REFERENCE TO RELATED APPLICATION

The present application is a continuation in part of USSN 08/813,507, filed March 7, 1997, which derives priority from provisional application 60/032,069, filed December 2, 1996, both of which are incorporated by reference in its entirety for all purposes.

BACKGROUND OF THE INVENTION

The genomes of all organisms undergo spontaneous mutation in the course of their continuing evolution generating variant forms of progenitor sequences (Gusella, Ann. Rev. Biochem. 55, 831-854 (1986)). The variant form may confer an evolutionary advantage or disadvantage relative to a progenitor form or may be neutral. In some instances, a variant form confers a lethal disadvantage and is not transmitted to subsequent generations of the organism. In other instances, a variant form confers an evolutionary advantage to the species and is eventually incorporated into the DNA of many or most members of the species and effectively becomes the progenitor form. In many instances, both progenitor and variant form(s) survive and co-exist in a species population. The coexistence of multiple forms of a sequence gives rise to polymorphisms.

Several different types of polymorphism have been reported. A restriction fragment length polymorphism (RFLP) means a variation in DNA sequence that alters the length of a restriction fragment as described in Botstein et al., Am. J. Hum. Genet. 32, 314-331 (1980). The restriction fragment length polymorphism may create or delete a restriction site, thus changing the length of the restriction fragment. RFLPs have been widely used in human and animal genetic analyses (see WO 90/13668; WO90/11369; Donis-Keller, Cell 51, 319-337 (1987); Lander et al., Genetics 121, 85-99 (1989)). When a heritable trait can be linked to a particular RFLP, the presence of the RFLP in an individual can be used to predict the likelihood that the animal will also exhibit the trait.

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Other polymorphisms take the form of short tandem repeats (STRs) that include tandem di-, tri- and tetra-nucleotide repeated motifs. These tandem repeats are also referred to as variable number tandem repeat (VNTR) polymorphisms. VNTRs have been used in identity and paternity analysis (US 5,075,217; Armour et al., FEBS Lett. 307, 113-115 (1992); Horn et al., WO 91/14003; Jeffreys, EP 370,719), and in a large number of genetic mapping studies.

Other polymorphisms take the form of single nucleotide variations between individuals of the same species. polymorphisms are far more frequent than RFLPs, STRs and VNTRs. Some single nucleotide polymorphisms occur in proteincoding sequences, in which case, one of the polymorphic forms may give rise to the expression of a defective or other variant protein. Other single nucleotide polymorphisms occur in noncoding regions. Some of these polymorphisms may also result in defective or variant protein expression (e.g., as a result of defective splicing). Other single nucleotide polymorphisms have no phenotypic effects. Single nucleotide polymorphisms can be used in the same manner as RFLPs, and VNTRs but offer several advantages. Single nucleotide polymorphisms occur with greater frequency and are spaced more uniformly throughout the genome than other forms of The greater frequency and uniformity of single polymorphism. nucleotide polymorphisms means that there is a greater

probability that such a polymorphism will be found in close proximity to a genetic locus of interest than would be the case for other polymorphisms. Also, the different forms of characterized single nucleotide polymorphisms are often easier to distinguish that other types of polymorphism (e.g., by use of assays employing allele-specific hybridization probes or primers).

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Despite the increased amount of nucleotide sequence data being generated in recent years, only a minute proportion of the total repository of polymorphisms has so far been identified. The paucity of polymorphisms hitherto identified is due to the large amount of work required for their detection by conventional methods. For example, a conventional approach to identifying polymorphisms might be to sequence the same stretch of oligonucleotides in a population of individuals by didoxy sequencing. In this type of approach, the amount of work increases in proportion to both the length of sequence and the number of individuals in a population and becomes impractical for large stretches of DNA or large numbers of subjects.

SUMMARY OF THE INVENTION

The invention provides nucleic acid segments containing at least 10, 15 or 20 contiguous bases from a fragment shown in Table 1 or 2, the complement thereof, or a plant cognate variant of the fragment. The segments include a polymorphic site shown in Table 1 or 2. The segments can be DNA or RNA, and can be double- or single-stranded. Some segments are 10-20 or 10-50 bases long. Preferred segments include a diallelic polymorphic site.

The invention further provides allele-specific oligonucleotides that hybridizes to a segment of a fragment shown in Table 1 or 2 or its complement. These oligonucleotides can be probes or primers. Also provided are isolated nucleic acids comprising a sequence of Table 1 or 2 or the complement thereto, in which the polymorphic site within the sequence is occupied by a base other than the reference base shown in Table 1 or 2.

The invention further provides a method of analyzing a nucleic acid from a subject. The method determines which base or bases is/are present at any one of the polymorphic sites shown in Table 1 or 2. Optionally, a set of bases occupying a set of the polymorphic sites shown in Table 1 or 2 is determined. This type of analysis can be performed on a plurality of subjects who are tested for the presence of a phenotype. The presence or absence of phenotype can then be correlated with a base or set of bases present at the polymorphic sites in the subjects tested.

BRIEF DESCRIPTION OF THE FIGURE

Fig. 1 shows probe arrays tiles for two allelic forms of the Brassica 18A2 polymorphism.

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DEFINITIONS

A nucleic acid, such an oligonucleotide, oligonucleotide can be DNA or RNA, and single- or double-stranded. Oligonucleotides can be naturally occurring or synthetic, but are typically prepared by synthetic means. Preferred nucleic acids of the invention include segments of DNA, or their complements including any one of the polymorphic sites shown in Table 1 or 2. The segments are usually between 5 and 100 bases, and often between 5-10, 5-20, 10-20, 10-50, 20-50 or 20-100 bases. The polymorphic site can occur within any position of the segment. The segments can be from any of the allelic forms of DNA shown in Table 1 or 2. Methods of synthesizing oligonucleotides are found in, for example, Oligonucleotide Synthesis: A Practical Approach (Gait, ed., IRL Press, Oxford, 1984).

Hybridization probes are oligonucleotides capable of binding in a base-specific manner to a complementary strand of nucleic acid. Such probes include peptide nucleic acids, as described in Nielsen et al., Science 254, 1497-1500 (1991).

The term primer refers to a single-stranded oligonucleotide capable of acting as a point of initiation of template-directed DNA synthesis under appropriate conditions (i.e., in the presence of four different nucleoside

triphosphates and an agent for polymerization, such as, DNA or RNA polymerase or reverse transcriptase) in an appropriate buffer and at a suitable temperature. The appropriate length of a primer depends on the intended use of the primer but typically ranges from 15 to 30 nucleotides. Short primer molecules generally require cooler temperatures to form sufficiently stable hybrid complexes with the template. A primer need not reflect the exact sequence of the template but must be sufficiently complementary to hybridize with a template. The term primer site refers to the area of the target DNA to which a primer hybridizes. The term primer pair means a set of primers including a 5' upstream primer that hybridizes with the 5' end of the DNA sequence to be amplified and a 3', downstream primer that hybridizes with the complement of the 3' end of the sequence to be amplified.

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Linkage describes the tendency of genes, alleles, loci or genetic markers to be inherited together as a result of their location on the same chromosome, and can be measured by percent recombination between the two genes, alleles, loci or genetic markers.

Polymorphism refers to the occurrence of two or more genetically determined alternative sequences or alleles in a population. A polymorphic marker or site is the locus at which divergence occurs. Preferred markers have at least two alleles, each occurring at frequency of greater than 1%, and more preferably greater than 10% or 20% of a selected population. A polymorphic locus may be as small as one base Polymorphic markers include restriction fragment length polymorphisms, variable number of tandem repeats (VNTR's), hypervariable regions, minisatellites, dinucleotide repeats, trinucleotide repeats, tetranucleotide repeats, simple sequence repeats, and insertion elements such as Alu. The first identified allelic form is arbitrarily designated as a the reference form and other allelic forms are designated as alternative or variant alleles. The allelic form occurring most frequently in a selected population is sometimes referred to as the wildtype form. Diploid organisms may be homozygous

or heterozygous for allelic forms. A diallelic polymorphism has two forms. A triallelic polymorphism has three forms.

A single nucleotide polymorphism occurs at a polymorphic site occupied by a single nucleotide, which is the site of variation between allelic sequences. The site is usually preceded by and followed by highly conserved sequences of the allele (e.g., sequences that vary in less than 1/100 or 1/1000 members of the populations).

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A single nucleotide polymorphism usually arises due to substitution of one nucleotide for another at the polymorphic site. A transition is the replacement of one purine by another purine or one pyrimidine by another pyrimidine. A transversion is the replacement of a purine by a pyrimidine or vice versa. Single nucleotide polymorphisms can also arise from a deletion of a nucleotide or an insertion of a nucleotide relative to a reference allele.

Hybridizations are usually performed under stringent conditions, for example, at a salt concentration of no more than 1 M and a temperature of at least 25°C. For example, conditions of 5X SSPE (750 mM NaCl, 50 mM NaPhosphate, 5 mM EDTA, pH 7.4) and a temperature of 25-30°C are suitable for allele-specific probe hybridizations.

The invention provides DNA sequences from Brassica and corn and cognate sequences from other plants. cognate refers to a gene sequence that is evolutionarily and functionally related between species. Cognate genes typically exhibit a high degree of sequence identity, (e.g., at least 80, 90, 95 or 99%) and hybridize to each other under stringent conditions. A polymorphic site in cognate sequences occur in corresponding positions when the sequences are maximally aligned according to the criteria of any one of the following references: Smith & Waterman (1981), Adv. Appl. Math. 2, 482; Needleman & Wunsch (1970), J. Mol. Biol. 48: 443; Pearson & Lipman (1988), Proc. Natl. Acad. Sci. (U.S.A.) 85: GAP, BESTFIT, FASTA, and TFASTA in the Wisconsin Genetics Software Package, Genetics Computer Group, 575 Science Dr., Madison, WI (each of which is incorporated by reference in its entirety for all purposes).

Nucleic acids of the invention are often in isolated form. An isolated nucleic acid means an object species that is the predominant species present (i.e., on a molar basis it is more abundant than any other individual species in the composition). Preferably, an isolated nucleic acid comprises at least about 50, 80 or 90 percent (on a molar basis) of all macromolecular species present. Most preferably, the object species is purified to essential homogeneity (contaminant species cannot be detected in the composition by conventional detection methods).

DESCRIPTION OF THE PRESENT INVENTION

I. Novel Polymorphisms of the Invention

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The present application provides oligonucleotides containing polymorphic sites isolated from two Brassica species, B. napus and B. oleracea, or corn, or cognate sequences from other plants. The invention also includes various methods for using those novel oligonucleotides to identify, distinguish, and determine the relatedness of individual strains or pools of nucleic acids from plants, particularly vegetal plants, and especially plants, within the family Cruciferae or Teosinte.

The genus Brassica is part of the family Cruciferae.

Members of the Brassica genus have been described as Old World

Temperate Zone herbs of the mustard family with beaked

cylindrical pods. Merriam-Webster's Collegiate Dictionary,

(10th ed., 1993), p.139. Many cruciferous plants are

important agricultural items and include many foodstuffs

(condiments, oilseeds, and vegetables). For example, canola

(a type of Brassica napus) is one of the largest crops in

Canada.

Sequences 1-149 in Table 1 and sequences from 86-5B5-N3 to 2B7/2F7-2A in Table 2 were isolated from B. napus and B. oleracea using oligonucleotide primers designed from expressed DNA sequences from Arabidopsis thaliana, a relative of Brassica napus and member of the Cruciferae family. See Hofte et al., An inventory of expressed sequence tags obtained by partial sequencing of cDNAs from Arabidopsis thaliana, Plant

J., Vol.4, pp. 1051-1061 (1993) and Newman et al., Genes Galore: A Summary of Methods for Accessing Results from Large-Scale Partial Sequencing of Anonymous Arabidopsis cDNA Clones, Plant Physiol., Vol. 106, pp. 1241-1255 (1994). There is a high degree of homology between the coding sequences of Arabidopsis, Brassica, and other members of the Cruciferae family.

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The designations in Table 1 are as follows. The first column is an arbitrarily assigned identification number for a polymorphism. The second and third columns, xmin and ymin, are the co-ordinates of probes for analyzing the polymorphism on a DNA chip (to be described below). The first number in the marker name is the Brassica strain name corresponding to the upper allele sequence. The next number designates the primer pair used for the PCR amplification. The sequences of some primers are described at the web site (http://www.yorku.ca/ftp/york other/cgat/) (incorporated by reference in its entirety for all purposes). The last number is the name of the strain for the lower allele sequence. example 85/5B5/86-1 means that the polymorphic site was identified by comparing strains 85 and 86-1 at a segment amplified by primers 5B5. Each sequence in the table includes a polymorphic site shown in square brackets [] and flanking bases common to both strains being compared. The upper and lower sequences in the square brackets are from the two strains being compared (upper strand corresponding to the first designated strain). A "/" within square brackets followed or preceded by a blank space represents an addition/deletion polymorphism. An asterisk indicates triallelic markers. The designation N indicates a base whose identity was not determined. The nomenclature for Table 2 is similar except that only fragment names and sequences flanking and including polymorphic sites are shown. In both Tables 1 and 2, the symbol "T" should be read as a "U" in RNA forms of the nucleic acids provided by the invention.

The invention further provides oligonucleotides from corn (zea maize), and cognate sequences from other plants especially those from the *Teosinte* family in which corn falls.

DNA was extracted from maize lines as described in Rogers and Bendich, Plant Biol. Manual A6, 1-10 (1988) with modification described in Murigneux et al., Theo. Appl. Genet. 86, 837-842 1993). PCR amplification was done on six maize lines representing a wide range of genetic variability and including both European flint material and US dent germplasm. maize lines were chosen to maximize genetic variability and thereby improve the probability of finding polymorphisms in the allelic sequences. Preferred primers for amplification 10 of some DNA segments including polymorphic sites are shown in Table 3. Other primers can be designed to incorporate sequences flanking the nucleic acid segments listed in Tables 1 and 2. Flanking sequences can be obtained from GenBank, or Bae et al., Maydica 35, 317-322 (1990), or PCR-based gene walking. See Parker et al., Nucl. Acids Res. 19, 3055-3060. 15 A specific primer for the sequenced region is primed with a general primer that hybridizes to the flanking region.

Nucleic acid segments flanking polymorphic sites in Zea Maize are shown as sequences 150-264 of Table 1, and sequences from S71G2/G6-1 to S01G3/G7-2 in Table 2. The nomenclature is the same as that described earlier for *Brassica sequences*.

II. Analysis of Polymorphisms

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A. <u>Preparation of Samples</u>

Polymorphisms are detected in a target nucleic acid from a plant being analyzed. Target nucleic acids can be genomic or cDNA. Many of the methods described below require amplification of DNA from target samples. This can be accomplished by e.g., PCR. See generally PCR Technology: Principles and Applications for DNA Amplification (ed. H.A. Erlich, Freeman Press, NY, NY, 1992); PCR Protocols: A Guide to Methods and Applications (eds. Innis, et al., Academic Press, San Diego, CA, 1990); Mattila et al., Nucleic Acids Res. 19, 4967 (1991); Eckert et al., PCR Methods and Applications 1, 17 (1991); PCR (eds. McPherson et al., IRL Press, Oxford); and U.S. Patent 4,683,202 (each of which is incorporated by reference for all purposes).

Other suitable amplification methods include the ligase chain reaction (LCR) (see Wu and Wallace, Genomics 4, 560 (1989), Landegren et al., Science 241, 1077 (1988), transcription amplification (Kwoh et al., Proc. Natl. Acad. Sci. USA 86, 1173 (1989)), and self-sustained sequence replication (Guatelli et al., Proc. Nat. Acad. Sci. USA, 87, 1874 (1990)) and nucleic acid based sequence amplification (NASBA). The latter two amplification methods involve isothermal reactions based on isothermal transcription, which produce both single stranded RNA (ssRNA) and double stranded DNA (dsDNA) as the amplification products in a ratio of about 30 or 100 to 1, respectively.

B. <u>Detection of Polymorphisms in Target DNA</u>

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There are two distinct types of analysis depending whether a polymorphism in question has already been characterized. The first type of analysis is sometimes referred to as de novo characterization. This analysis compares target sequences in different individual plants to identify points of variation, i.e., polymorphic sites. The de novo identification of the polymorphisms of the invention is described in the Examples section. The second type of analysis is determining which form(s) of a characterized polymorphism are present in plants under test. There are a variety of suitable procedures, which are discussed in turn.

1. Allele-Specific Probes

The design and use of allele-specific probes for analyzing polymorphisms is described by e.g., Saiki et al., Nature 324, 163-166 (1986); Dattagupta, EP 235,726, Saiki, WO 89/11548. Allele-specific probes can be designed that hybridize to a segment of target DNA from one member of a species but do not hybridize to the corresponding segment from another member due to the presence of different polymorphic forms in the respective segments from the two members. Hybridization conditions should be sufficiently stringent that there is a significant difference in hybridization intensity between alleles, and preferably an essentially binary

response, whereby a probe hybridizes to only one of the alleles. Some probes are designed to hybridize to a segment of target DNA such that the polymorphic site aligns with a central position (e.g., in a 15 mer at the 7 position; in a 16 mer, at either the 8 or 9 position) of the probe. This design of probe achieves good discrimination in hybridization between different allelic forms.

Allele-specific probes are often used in pairs, one member of a pair showing a perfect match to a reference form of a target sequence and the other member showing a perfect match to a variant form. Several pairs of probes can then be immobilized on the same support for simultaneous analysis of multiple polymorphisms within the same target sequence.

2. Tiling Arrays

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The polymorphisms can also be identified by hybridization to nucleic acid arrays, some example of which are described by WO 95/11995 (incorporated by reference in its entirety for all purposes). One form of such arrays is described in the Examples section in connection with de novo identification of polymorphisms. The same array or a different array can be used for analysis of characterized polymorphisms. WO 95/11995 also describes subarrays that are optimized for detection of a variant forms of a precharacterized polymorphism. Such a subarray contains probes designed to be complementary to a second reference sequence, which is an allelic variant of the first reference sequence. The second group of probes is designed by the same principles as described in the Examples except that the probes exhibit complementarity to the second reference sequence. inclusion of a second group (or further groups) can be particular useful for analyzing short subsequences of the primary reference sequence in which multiple mutations are expected to occur within a short distance commensurate with the length of the probes (i.e., two or more mutations within 9 to 21 bases).

A DNA chip has been designed containing subarrays of probes for analyzing each of the 264 polymorphic sites shown

in Table 1. Preferred polymorphic sites for inclusion on such a chip are selected by a number of criteria. Some polymorphic sites are selected because they are evenly distributed throughout the genome of a plant. For example, one or more polymorphisms is present on each chromosome. Preferably, the polymorphisms are no more than 20 cM apart. Other polymorphisms are selected because they occur within a segment that can be made to give a single band on Southern analysis. Other polymorphisms may be selected because they occur in genes encoding enzymes that function in the same pathway.

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Preferred primer pairs for amplification of nucleic acid segments flanking most of these polymorphic sites of Table 1 are shown in Table 3. This chip is used to determine a polymorphic profile of a plant under test at some or all of the polymorphic sites on the chip. Thus, one can determine which allelic form is present in a plant at up to 264 sites. Such a profile is useful in some of the applications described below.

DNA segments are preferably amplified in a multiplex For example, polymorphic sites 5, 8, 64, 108, 36, 38, and 136 (Table 1) can be amplified from the primers shown in Table 3 in one group. Polymorphic sites 11, 12, 15, 29, 130, 139, 140, 40, 49, 63, 72, 81, 86, 92, 93, 102, 142, 146, 149, 16, 138 can be amplified in another group. Polymorphic sites 1, 7, 17, 25, 26, 42, 60, 67, 73, 75, 106, 107, 111, 123, 133, 43, 51, 82, 95, and 101 can be amplified in another group. Polymorphic sites 2, 9, 20, 34, 41, 45, 61, 62, 65, 74, 75, 80, 91, 100, 105, 110, 123, and 14 can be amplified in another group. Polymorphic sites 4, 41, 61, 62, 65, 66, 76, 80, 91, 94, 111, 114, 121, 132, 22, 32, 33, 44, 53, 97 and 145 can be amplified in another group. Polymorphic sites 6, 21, 27, 46, 66, 68, 71, 77, 89, 89, 98, 103, 105, 31, 35, 54, 59, 84, 85, 96, 116, 137, 141, and 148 can be amplified in another Polymorphic sites 48, 70, 78, 87, 88, 90, 99, 103, 104, 112, 117, 118, 119, 122, 124, 13, 18, 30, 37, 69, 94, 135, and 147 can be amplified in another group. Polymorphic sites 247, 207, 203, and 173 can be amplified in another group. Polymorphic sites 229, 217, 212, 204, 195, 171, 175,

236, 174, and 193 can be amplified in another group.
Polymorphic sites 232, 243, 157, 162, 169, 177, 179, 234, 241, 255, 152, 160, 184, 188, 192, 199, 201, 205, 210, 220, 223, 226, 245, 249, and 252 can be amplified in another group.

Polymorphic sites 233, 244, 246, 248, 227, 211, 214, 200, 150, 158, 170, 178, 181, 238, 254, 161, 165, 180, 187, 222, and 231 can be amplified from another group. Polymorphic sites 228, 235, 215, 218, 221, 225, 198, 155, 163, 166, 167, 172, 216, 242, 159, 182, 186, 191, 202, 208, 230, 237, and 253 can be amplified in another group. Polymorphic sites 219, 224, 206, 209, 213, 194, 196, 197, 151, 153, 156, 164, 185, 250, 154, 168, 176, 183, 190, 239, 251, and 256 can be amplified in another group.

3. Allele-Specific Primers

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An allele-specific primer hybridizes to a site on target DNA overlapping a polymorphism and only primes amplification of an allelic form to which the primer exhibits perfect complementarity. See Gibbs, Nucleic Acid Res. 17, 2427-2448 (1989). This primer is used in conjunction with a second primer which hybridizes at a distal site. Amplification proceeds from the two primers leading to a detectable product signifying the particular allelic form is present. A control is usually performed with a second pair of primers, one of which shows a single base mismatch at the polymorphic site and the other of which exhibits perfect complementarity to a distal site. The single-base mismatch prevents amplification and no detectable product is formed. The method works best when the mismatch is included in the 3'most position of the oligonucleotide aligned with the polymorphism because this position is most destabilizing to elongation from the primer. See, e.g., WO 93/22456.

4. <u>Direct-Sequencing</u>

The direct analysis of the sequence of polymorphisms of the present invention can be accomplished using either the dideoxy chain termination method or the Maxam Gilbert method (see Sambrook et al., Molecular Cloning, A

Laboratory Manual (2nd Ed., CSHP, New York 1989); Zyskind et al., Recombinant DNA Laboratory Manual, (Acad. Press, 1988)).

- Amplification products generated using the polymerase chain reaction can be analyzed by the use of denaturing gradient gel electrophoresis. Different alleles can be identified based on the different sequence-dependent melting properties and electrophoretic migration of DNA in solution. Erlich, ed., PCR Technology, Principles and Applications for DNA Amplification, (W.H. Freeman and Co, New York, 1992), Chapter 7.
- Single-Strand Conformation Polymorphism Analysis 6. Alleles of target sequences can be differentiated 15 using single-strand conformation polymorphism analysis, which identifies base differences by alteration in electrophoretic migration of single stranded PCR products, as described in Orita et al., Proc. Nat. Acad. Sci. 86, 2766-2770 (1989). Amplified PCR products can be generated as described above, 20 and heated or otherwise denatured, to form single stranded amplification products. Single-stranded nucleic acids may refold or form secondary structures which are partially dependent on the base sequence. The different electrophoretic mobilities of single-stranded amplification products can be 25 related to base-sequence difference between alleles of target sequences.

III. Methods of Use

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After determining polymorphic form(s) present in a subject plant at one or more polymorphic sites, this information can be used in a number of methods.

A. Fingerprint Analysis

Analysis of which polymorphisms are present in a plant is useful in determining of which strain the plant is a member an in distinguishing one strain from another. A genetic fingerprint for an individual strain can be made by determining the nucleic acid sequence possessed by that

individual strain that corresponds to a region of the genome known to contain polymorphisms. For a discussion of genetic fingerprinting in the animal kingdom, see, for example, Stokening et.al., Am. J. Hum. Genet. 48:370-382 (1991). The probability that one or more polymorphisms in an individual strain is the same as that in any other individual strain decreases as the number of polymorphic sites is increased.

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The comparison of the nucleic acid sequences from two strains at one or multiple polymorphic sites can also demonstrate common or disparate ancestry. Since the polymorphic sites are within a large region in the genome, the probability of recombination between these polymorphic sites is low. That low probability means the haplotype (the set of all the disclosed polymorphic sites) set forth in this application should be inherited without change for at least several generations. Knowledge of plant strain or ancestry is useful, for example, in a plant breeding program or in tracing progeny of a proprietary plant. Fingerprints are also used to identify an individual strain and to distinguish or determine the relatedness of one individual strain to another. Genetic fingerprinting can also be useful in hybrid certification, the certification of seed lots, and the assertion of plant breeders rights under the laws of various countries.

Genetic fingerprinting is also useful in screening progeny of a backcross for a high contribution of a parent strain (see Hospital et al., Genetics 44, 843-874 (1992)). In a backcross, a progeny that has been bred to have a desired phenotype is crossed with a parental strain to remove genetic variations that do not contribute to the desired phenotype but which may have latent undesirable effects. After crossing, offspring retaining the desired phenotype are identified, and their polymorphic profile is determined. The offspring retaining the desired phenotype and having the greatest similarity in polymorphic profile to a parent strain can then used as a production model.

B. Correlation of Polymorphisms with Phenotypic Traits

The polymorphisms of the invention may contribute to the phenotype of a plant in different ways. Some polymorphisms occur within a protein coding sequence and contribute to phenotype by affecting protein structure. The effect may be neutral, beneficial or detrimental, or both beneficial and detrimental, depending on the circumstances. Other polymorphisms occur in noncoding regions but may exert phenotypic effects indirectly via influence on replication, transcription, and translation. A single polymorphism may affect more than one phenotypic trait. Likewise, a single phenotypic trait may be affected by polymorphisms in different genes. Further, some polymorphisms predispose a plant to a distinct mutation that is causally related to a certain phenotype.

Phenotypic traits include characteristics such as growth rate, crop yield, crop quality, resistance to pathogens, herbicides, and other toxins, nutrient requirements, resistance to high temperature, freezing, drought, requirements for light and soil type, aesthetics, and height. Other phenotypic traits include susceptibility or resistance to diseases, such as plant cancers. Often polymorphisms occurring within the same gene correlate with the same phenotype.

Correlation is performed for a population of plants, which have been tested for the presence or absence of a phenotypic trait of interest and for polymorphic markers sets. To perform such analysis, the presence or absence of a set of polymorphisms (i.e. a polymorphic set) is determined for a set of the plants, some of whom exhibit a particular trait, and some of which exhibit lack of the trait. The alleles of each polymorphism of the set are then reviewed to determine whether the presence or absence of a particular allele is associated with the trait of interest. Correlation can be performed by standard statistical methods such as a κ -squared test and statistically significant correlations between polymorphic form(s) and phenotypic characteristics are noted.

Correlations between characteristics and phenotype are useful for breeding for desired characteristics. By analogy, Beitz et al., US 5,292,639 discuss use of bovine mitochondrial polymorphisms in a breeding program to improve milk production in cows. To evaluate the effect of mtDNA D-loop sequence polymorphism on milk production, each cow was assigned a value of 1 if variant or 0 if wildtype with respect to a prototypical mitochondrial DNA sequence at each of 17 locations considered. Each production trait was analyzed individually with the following animal model:

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 $Y_{ijkpn} = \mu + YS_i + P_j + X_k + \beta_1 + \dots + \beta_{17} + PE_n + a_n + e_p$ where Yijkno is the milk, fat, fat percentage, SNF, SNF percentage, energy concentration, or lactation energy record; μ is an overall mean; YS $_{i}$ is the effect common to all cows calving in year-season; X_k is the effect common to cows in either the high or average selection line; β_1 to β_{17} are the binomial regressions of production record on mtDNA D-loop sequence polymorphisms; PEn is permanent environmental effect common to all records of cow n; an is effect of animal n and is composed of the additive genetic contribution of sire and dam breeding values and a Mendelian sampling effect; and en is a random residual. It was found that eleven of seventeen polymorphisms tested influenced at least one production trait. Bovines having the best polymorphic forms for milk production at these eleven loci are used as parents for breeding the next generation of the herd.

One can test at least several hundreds of markers simultaneously in order to identify those linked to a gene or chromosomal region. For example, to identify markers linked to a gene conferring disease resistance, a DNA pool is constructed from plants of a segregating population that are resistant and another pool is constructed from plants that are sensitive to the disease. Those two DNA pools are identical except for the DNA sequences at the resistance gene locus and in the surrounding genomic area. Hybridization of such DNA pools to the DNA sequences listed in Table 1 allows the simultaneous testing of several hundreds of loci for polymorphisms. Allelic polymorphism-detecting sequences that

show differences in hybridization patterns between such DNA pools represent loci linked to the disease resistance gene.

The method just described can also be applied to rapidly identify rare alleles in large populations of plants. For example, nucleic acid pools are constructed from several individuals of a large population. The nucleic acid pools are hybridized to nucleic acids having the polymorphism-detecting sequences listed in Table 1 or 2. The detection of a rare hybridization profile will indicate the presence of a rare allele in a specific nucleic acid pool. RNA pools are particularly suited to identify differences in gene expression.

IV. Modified Polypeptides and Gene Sequences

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The invention further provides variant forms of nucleic acids and corresponding proteins. The nucleic acids comprise at least 10 contiguous bases of one of the sequences described in Table 1 or 2, in any of the allelic forms shown. Some nucleic acid encode full-length proteins.

Genes can be expressed in an expression vector in which a gene is operably linked to a native or other promoter. Usually, the promoter is a eukaryotic promoter for expression in a eukaryotic cell. The transcription regulation sequences typically include a heterologous promoter and optionally an enhancer which is recognized by the host. The selection of an appropriate promoter, for example trp, lac, phage promoters, glycolytic enzyme promoters and tRNA promoters, depends on the host selected. Commercially available expression vectors can be used. Vectors can include host-recognized replication systems, amplifiable genes, selectable markers, host sequences useful for insertion into the host genome, and the like.

The means of introducing the expression construct into a host cell varies depending upon the particular construction and the target host. Suitable means include fusion, conjugation, transfection, transduction, electroporation or injection, as described in Sambrook, supra. A wide variety of host cells can be employed for expression of the variant gene, both prokaryotic and eukaryotic. Suitable host cells include

bacteria such as E. coli, yeast, filamentous fungi, insect cells, mammalian cells, typically immortalized, e.g., mouse, CHO, human and monkey cell lines and derivatives thereof, and plant cells. Preferred host cells are able to process the variant gene product to produce an appropriate mature polypeptide. Processing includes glycosylation, ubiquitination, disulfide bond formation, general posttranslational modification, and the like.

The DNA fragments are introduced into cultured plant 10 cells by standard methods including electroporation (From et al., Proc. Natl Acad. Sci. USA 82, 5824 (1985), infection by viral vectors such as cauliflower mosaic virus (CaMV) (Hohn et al., Molecular Biology of Plant Tumors, (Academic Press, New York, 1982) pp. 549-560; Howell, US 4,407,956), high velocity 15 ballistic penetration by small particles with the nucleic acid either within the matrix of small beads or particles, or on the surface (Klein et al., Nature 327, 70-73 (1987)), use of pollen as vector (WO 85/01856), or use of Agrobacterium tumefaciens transformed with a Ti plasmid in which DNA 20 fragments are cloned. The Ti plasmid is transmitted to plant cells upon infection by Agrobacterium tumefaciens, and is stably integrated into the plant genome (Horsch et al., Science, 233, 496-498 (1984); Fraley et al., Proc. Natl. Acad. Sci. USA 80, 4803 (1983)).

The protein may be isolated by conventional means of protein biochemistry and purification to obtain a substantially pure product, i.e., 80, 95 or 99% free of cell component contaminants, as described in Jacoby, Methods in Enzymology Volume 104, Academic Press, New York (1984);

Scopes, Protein Purification, Principles and Practice, 2nd Edition, Springer-Verlag, New York (1987); and Deutscher (ed), Guide to Protein Purification, Methods in Enzymology, Vol. 182 (1990). If the protein is secreted, it can be isolated from the supernatant in which the host cell is grown. If not secreted, the protein can be isolated from a lysate of the host cells.

The invention further provides transgenic plants capable of expressing an exogenous variant gene and/or having

one or both alleles of an endogenous variant gene inactivated. Plant regeneration from cultural protoplasts is described in Evans et al., "Protoplasts Isolation and Culture," Handbook of Plant Cell Cultures 1, 124-176 (MacMillan Publishing Co., New 5 York, 1983); Davey, "Recent Developments in the Culture and Regeneration of Plant Protoplasts, " Protoplasts, (1983) - pp. 12-29, (Birkhauser, Basal 1983); Dale, "Protoplast Culture and Plant Regeneration of Cereals and Other Recalcitrant Crops," Protoplasts (1983) - pp. 31-41, (Birkhauser, Basel 1983); Binding, "Regeneration of Plants," Plant Protoplasts, pp. 21-10 73, (CRC Press, Boca Raton, 1985). For example, a variant gene responsible for a disease-resistant phenotype can be introduced into the plant to simulate that phenotype. Expression of an exogenous variant gene is usually achieved by 15 operably linking the gene to a promoter and optionally an enhancer. Inactivation of endogenous variant genes can be achieved by forming a transgene in which a cloned variant gene is inactivated by insertion of a positive selection marker. See Capecchi, Science 244, 1288-1292 (1989). Such transgenic 20 plants are useful in a variety of screening assays. For example, the transgenic plant can then be treated with compounds of interest and the effect of those compounds on the disease resistance can be monitored. In another example, the transgenic plant can be exposed to a variety of environmental conditions to determine the effect of those conditions on the 25 resistance to the disease.

In addition to substantially full-length polypeptides, the present invention includes biologically active fragments of the polypeptides, or analogs thereof, including organic molecules which simulate the interactions of the peptides. Biologically active fragments include any portion of the full-length polypeptide which confers a biological function on the variant gene product, including ligand binding, and antibody binding. Ligand binding includes binding by nucleic acids, proteins or polypeptides, small biologically active molecules, or large cellular structures.

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Polyclonal and/or monoclonal antibodies that specifically bind to one allelic gene products but not to a

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second allelic gene product are also provided. Antibodies can be made by injecting mice or other animals with the variant gene product or synthetic peptide fragments thereof. Monoclonal antibodies are screened as are described, for 5 example, in Harlow & Lane, Antibodies, A Laboratory Manual, Cold Spring Harbor Press, New York (1988); Goding, Monoclonal antibodies, Principles and Practice (2d ed.) Academic Press, New York (1986). Monoclonal antibodies are tested for specific immunoreactivity with a variant gene product and lack of immunoreactivity to the corresponding prototypical gene product. These antibodies are useful in diagnostic assays for detection of the variant form, or as an active ingredient in a pharmaceutical composition.

15 V. <u>Kits</u>

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The invention further provides kits comprising at least one allele-specific oligonucleotide as described above. Often, the kits contain one or more pairs of allele-specific oligonucleotides hybridizing to different forms of a polymorphism. In some kits, the allele-specific oligonucleotides are provided immobilized to a substrate. example, the same substrate can comprise allele-specific oligonucleotide probes for detecting at least 10, 100 or all of the polymorphisms shown in Table 1 or 2. Optional additional components of the kit include, for example, restriction enzymes, reverse-transcriptase or polymerase, the substrate nucleoside triphosphates, means used to label (for example, an avidin-enzyme conjugate and enzyme substrate and chromogen if the label is biotin), and the appropriate buffers for reverse transcription, PCR, or hybridization reactions. Usually, the kit also contains instructions for carrying out the methods.

EXAMPLES

35 As noted above, the sequences in Table 1 were isolated from B. napus and B. oleracea using oligonucleotide primers designed from expressed DNA sequences from Arabidopsis thaliana, a relative of Brassica napus and member of the

Cruciferae family. Primers used to amplify B. napus and B. oleracea alleles were selected for an optimal length of 20 bases ± 2 based such that their melting temperatures were between 60°C and 65°C. Primers were synthesized on a 20 nmole scale using a high throughput DNA synthesizer capable of producing 96 primers simultaneously in a 96-well format. See Lashkari et al., Proc. Nat. Acad. Sci. 92, 7912-7915 (1995). The primers, which have an average length of 21 bases, were positioned within DNA sequences such that PCR products produced with cDNA templates would range between 100 and 450 bp. As introns in Arabidopsis genes are of modest size, 60% of the 1,920 primers tested on plant DNA gave PCR products.

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The components needed for PCR amplification were mixed in the following proportions for a 96 well microamp tray 206:1 of 10X PCR reaction buffer, 206:1 of 2 mM dNTPs, 186:1 of 15 mM MgCl $_2$, 720:1 of sterile ddH $_2$ 0 and, 20:1 of Taq DNA polymerase (Perkin Elmer). The enzyme was added just prior to dispensing 168:1 of this master mix into 8 20:1 of the appropriate forward and reverse primer 10 pmol/l stock solutions was added to each tube. A volume of 14:1 of this mixture was dispensed into each well of the microamp assembly with a BioHit 8-channel pipette. A volume of 5:1 of 20 ng/l template DNA solutions was added to the microamp assembly with a 12-channel pipette. The assembly was centrifuged for 30 sec to ensure that all reagents were mixed. Amplifications were performed in a Perkin Elmer system 9600 thermal cycler with an initial denaturation at 95°C for 1 min followed by 40 cycles of 94°C for 30 sec, 55°C for 30 sec, 72°C for 30 sec and a final extension at 72°C for 5 min. Products were separated by electrophoresis at 120 volts for 1 hr through 2% (w/v) agarose gels prestained with ethidium bromide. The banding patterns of these gels were recorded with an Alpha Innotech gel documentation system.

Any two amplicons obtained from the same primer set with two different plant varieties are said to be homomorphic if they have the same size. A set of 355 homomorphic Brassica napus and 250 homomorphic Brassica oleracae fragments were purified with Quiaquick columns and sequenced using dye

labeled dideoxy-terminators. See Stryer, Biochemistry 2nd. ed., pp. 592-593 (1981). The same primers used for the PCR amplification of the homomorphic DNA fragments were also used for the DNA sequencing of these fragments. The sequences obtained were aligned to identify single nucleotide polymorphisms.

Using VLSIPSTM technology (US 5,143,854; WO 90/15070; WO 92/10092), GeneChipJ was constructed using 20mer-probe sets to identify by hybridization the presence or absence of many 10 of the polymorphisms shown in Table 1 in a sample of plant nucleic acid. The tiling strategy used to create the GeneChipJ is set forth in Figure 1. Tiling strategies can be devised using the guidance provided herein by those skilled in the art. Tiling arrays are described in PCT/US94/12305 15 (incorporated by reference in its entirety for all purposes). ATiling@ generally means the synthesis of a defined set of oligonucleotide probes that is made up of a sequence complementary to the sequence to be analyzed (the target sequence), as well as preselected variations of that sequence. The variations usually include substitution at one or more 20 base positions with one or more nucleotides. Tiling strategies are discussed in Published PCT Application No. WO 95/11995 (incorporated by reference in its entirety for all purposes). In general, with a tiled array containing 4L 25 probes one can query every position in a nucleotide containing L number of bases. A 4L tiled array, for example, contains L number of sets of 4 probes, i.e. 4L probes. Each set of 4 probes contains the perfect complement to a portion of the target sequence with a single substitution for each nucleotide 30 at the same position in the probe. See also Chee, M., et. al., Science, October, 1996.

The tiling strategy for 20mer probes shown in Figure 1 for a single allele of the polymorphism employed probe sets having a perfect match and a corresponding single-base mismatch at the tenth base in the probe, counting from the 3= end. Each set had 14 pairs of probes that began at 14 successively shifted positions such that the substituted base lay from 7 bases upstream to 6 bases downstream from the

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polymorphic site. Two such sets of 28 probes were included to query the polymorphic site for the two alleles, as shown for example, in Figure 1. This collection of 56 probes constituted a detection block. Two such blocks per marker were synthesized to query both the forward and reverse strands. Thus each marker interrogated by the GeneChipJ was represented by a full set of 112 probes.

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All publications and patent applications cited above are incorporated by reference in their entirety for all purposes to the same extent as if each individual publication or patent application were specifically and individually indicated to be so incorporated by reference. Although the present invention has been described in some detail by way of illustration and example for purposes of clarity and understanding, it will be apparent that certain changes and modifications may be practiced within the scope of the appended claims.

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erassic	A POL	YMORE	HIC MARKERS	
NUMBER	XMIN	YMIN	MARKER_NAME	SEQUENCE
1	0	3	85/5B5/86-1	AGCAAGCTTACATGCGTGGA (GT/AA) GAGAGTCCTCGAGATCAACC
2	33	3	85/5B12/N3-1	CCTTGATCTCTCAAGTAATC[A/G]TCTCACCGGAAGATCCCTGA
3	66	3	85/5C3/86-2	ACCATCCATTAAACTGTATC[A/G]TCGCAATCTAACCAAAAGTT
4	99	3	85/5E1/86-1	TAAAGCAAAGAGAGTCTTAC [C/A]GTCTGCTGCATGATATACCC
5	132	3	85/5E1/86-2	CTACTGATAGTGAACCACCC (A/C)ATCCCCAAATTTAAAGCAAA
6	165	3	85/6A11/86	ATCCTATTGGTAGTAACACA [G/A]ATTGAGTTAATGTTGCAGGG
7	198	3	N1/6A11/N2	AGGCAAAGCGGTAGTTGCAA (G/A) ACTGCTTCTCACGAGGTAAT
8	231	3	N1/6A9/N2-1	CCAGCTTCAATGTCTGCATG[C/A]TTGTGTCGATGCCAAAGTTC
9	0	11	N1/6A9/N2-2	AAAGTTCATTACGATGATCT (A/G)ACCCTGCAGTCATCCATGGA
10	33	11	85/6A12/86	CTTCCCCCCCTCAATACCTC [T/G]TTCAAAAGTGAAAAGTGCAG
11	66	11	N1/6D1/N2-1	ATTTTGTTTTGTTCTGTC(G/C)GGTCAGGTCAGAACAAAGTT
12	99	11	N1/6H5/N2	AAACCAGAGCCACCTCCTTA(C/)CCACCTCATCGTTTCCTTTC
13	132	11	86/6F11/N2-2	GATTTCGACCGCAGTCTCAC (G/T)GAGGATGAGTATATCGCTTT
14	165	11	N1/6F11/N2	TAGGACAGGCAAACAATCTA(C/A)GCGGTCAAAATCCGATTTCG
15	198	11	L4/8A2/L6-2	AGCAGTGCAACAGCTCCTGC (A/T)AAGTCCCTGACGTACGAGGA
16	231	11	N1/8B5/N2	ACTCAAAAAACGATACCTC[G/C]GCCGTCTCTCGCCGTCTCCC
17	0	19	N1/8D4/N2-1	CAGGAGACAGTTACAGTCCC[/A]CAGAGTCGCAAGGATCTCGAA
18	33		85/8D4/86-2	CTGATCTTGAAGGAGACC[A/G]CCACAAGGTTCCATCCTATG
19	66	19	85/8H11/86	AGTGCGAGGCTCAGTTGGAT (G/T)ATTAGGGTGTCAGTAAATCA
20	99	19	85/10B8/86	NAGGTCCATGATGACAA (T/A) AAAGGTATTCCACATGTCAA
21	132	19	NZ/10B8/N3-2	ACATCCAACTTTTCTCCAGT[T/C]CTTTATTCTATCCTGATTTG
22	165		N2/10B8/N3-1	AAGGTATTCCATTGGTATAC (A/C)TCCAACTTTTCTCCAG1 TCT
23	198	19	85/10B9/86	GACCTTCTTGGGAAAGAAAG [T/C]TGTAACCGCGTCGAGATTCG
24	231	19	L5/10C8/N2	ATAGAAACCGCCGATGCTCA [G/A] GGACACGCCACCGTCTTCGT
25	0	27	L6/10C8/N2	CACTTTCTTCGTGGCTAAAT [G/T] CTTCGGCCGAGCCGGTCTCA
26	33		L6/10D2/N1	GTTATCATCAGTACCGGTAT [C/T]AACCCCAAGGCTAATTCTTA
27	66	27	85/10D2/L6	TTGGGTATCTACGGACTGAT[C/T]ATCGCTGTTATCATCAGTAC
28	99		N1/10E12/N2-1	GGAATTCAATACTCGCCAAC[G/T]TCTTCATTGCTGTCGTCGGC
29	132		N1/10E12/N2-2	TCCTTACGCCTTCAAGCGCA(C/G)CGCTCGCTCATGGGTGTCC
30	165	27	N1/10F4/N2	TGTATCTATGCGGTGGCTGC (G/C)GTCTCCGTTCGCGCCAGTAC
31	198	27	L1/10F4/N2*	GCGCCAGTACCGCCGGTTAC [C/A]ATCTCACTGCCTTCACGTCC
32	231	27	85/10F4/N2*	GCGCCAGTACCGCCGGTTAC (G/A) ATCTTAATGCCTTCACGTTC
33	0	35	85/10F9/N1-2	AACTTGGAATTCCACAACTT[G/C]AGAAACTTCGATGTGGTGCC
34	33	35	85/10F9/86	CGGTACTGCGAAAGCTGGAG[C/G]ATCAACTTGGAATTCCACAA
35	66	35	86/10F12/L3	AAAAGTGCTATTGTTCAGGT [G/C] GATGCTGCTCCGTTCAAGCA
36	99		85/10H6/86	GTCAAAAGCCACGGATTCAA (G/A) AACGTGCTCTTCTTGCGCCT
37	132		L1/10H6/L6	GCGGTGGZACAGGCGCTCA[G/T]TTTCTCGTCAAAAGCCACGG
38	165		85/10F12/86	AAACCAGGGTCCTTGATGTG [T/]GTCTACAACGCTTCCAACAA
39	198		85/11B7/86	
40	231			TTTGGGACCGTTCATGCCT[C/T]TGACCCATGTTCTTGCCACC
41	0	~~~	85/11C4/86 85/11D4/86-2	TTTTGGGACCGTTGGAGTTGC (A/G) TCTGCGGCTATGACGGTGGA
42	33		85/11D4/86-2	AATCTTTGCCATTGCTGTCA(A/G)TATCTTCGTCAGCTTCAGCT
43			L4/11A3/L1	TTGAAGGAGGTTGGTACACA[C/G]TTCTTCGAGCTACCGGAGAC
44	66 99	. ———	N2/11D11/N3	GACAACGCTGGTGCTATTGC C/T GAAATGGCTGGAATGAGCCA
**	וצב	43	86/11D11/N3	GCTGCTCTAGGGATGCTCAG [C/T] ACCATCGCCACCGGTTTGGC

45	132	43	85/11D11/86	ATGCTCAGCACCATCGCCAC [T/C]GGTTTGGCGATTGATGCTTA
-46	165	43	N2/11E3/N8a	GAGAAAGTGCTTGTGGAGAT (C/T)TACAAGTCCATACTGATGGC
47	198	43	86/11E3/N2a	AATGCTTGTGGAGATLTACA (G/A)GTCCATACTGATGGCGCAGG
48	231	43	86/11E3/N2b	AATGCTTGTGGAGATCTACA (G/A)GTCCATACTGATGGCGCAGG
49	0	51	85/11F12/86	AATGATTGGTTTGAGAAGCA [T/A] ACAGCTGGTACGCTTGATAT
50	33	51	85/11F7/86	GATAGGGCGAAGAGAGGGAA [G/A] AGTCCTGAGAGGAAAGAGAT
51	66	51	85/11H2/86-2	CTCTCTCCACAAAGACAC(A/C)GCTTTCTCCATGACCTTCGG
52	99	51	85/11H5/86-2	TCTCTGACGTCATGAAAGCT [C / A] ATGGCAAAATTGCTGATGGA
53	132	51	85/11H6/86-1a	GTTATCGATCGCGTGGTCCG[T/C]GAAACCCAAAATaCACCTTT
54	165	51	85/11H6/86-1b	GTTATCGATCGCGTGGTCCG[T/C]GAAACCCAAAATECACCTTT
55	198	51	85/12B6/N3	CGTCAGCCTTCTTCCGCCGC [A/C]GTCGTCCTCCGCAACCGTGC
56	231	51	86/12B6/85a	TGTCTCTTCCGTCAGCCTTC [C / T] TCCGCCGCAGTCGTCCTCCG
57	0	59	86/12B6/85b	TGTCTCTTCCGTCAGCCTTC[C/T]TCCGCCGCcGTCGTCCTCCG
58	33	59	86/12B11/85	TCAGGTTTACCTCTATATAT [T/] ATATTTCATGGTATGAAGGT
59	66	59	n1/12B11/N2-2	TATCCTGCAAATTGACATTT(T/C)CCTTCAGGTTCTAGAAGCTG
60	99	59	85/12C2/86	CGAGAACAGAAGAGAGAGA [C/]TGGAACACGTCGGACAGTAC
61	132	59	L3/12C7/L5	TGNCACAACGAAGGTTTTGG[C/T]GGAGGTAAATGCCGTGGGTT
62	165	59	L6/12C11/N2	ACCOGTCCTAGCGCCATGGC[C/T]ATTTTCCTCACCGTTTCTGG
63	198	59	N1/12D10/L6	TTGGGCTTTCGGTGGTATGA[T/]CTTCGTCCTCGTCTATTGCA
64	231	59	L2/12E10/L3	CACAAAGGTCTGCCTCACAA [G/T]TTCTACCACGGTCGTACTGG
65	0	67	85/12F4/86-1	TCCTTGATTCCTTAATAATC (A/T)TTGGCTGGGGGTCTTTCTAA
66	33	67	L1/12G5/N1	GCTTGAATAACGATGTCTAC (A/T) CTGCCTCGGCGTACGGCGGA
67	66	67	85/12G8/L1 ·	CTAAAAAGATCGACGAGTGT (C/T) CCTTACTACGCTCCATCTAT
68	99	67	L6/12G9/N1-1*	AGGTGGGTTTAGCGTGGCAT (T/C) CGATCCATTGGATGGATECA
69	132	67	85/12G9/L3*	NGTGGGTTTACCGTATCATT(T/C)GATCCATTGGATGGATCGAG
70	165	67	L3/12B11/N2-1	GCGGATCCTATATTGGGTCT[C/T]GATGGATTGTTTCTATCCCG
71	198	67	L2/12B11/N2-2	TATCCTGCAAATTGACATTT [T/C] CCTTCAGGTTCTAGAAGCTG
72	231	67	N1/12E10/L1	TACCACGGTCGTACTGGTCG[A/]TGTCTGGAACGTCACCAAGC
73	0	75	N1/13A3/N2a	CTGTCTCAgTTTGTTGGATC(C/G)AAATCGAAATCGAAAGCGTAC
74	33	75	N1/13A3/N2b	CTGTCTCAGTTTGTTGGATC(C/G)AAATC&AATCGAAAGCGTAC
75	66	75	L2/13E8/N2	ACACTGTTGGAGGACGTGAA (T/G) AAGATATTCAAGACAACATC
76.	99	75	N1/13F6/N2-2	TCTTTCGTATCTTGCTGAGT C/T GTTACGCCTGTCAACACCCG
77	132	75	L2/13F8/N2-1	GGAACCCTAGGGAGCCCACA(T/G)CTCCTTATGCTAAGCGGCGT
78	165	75	L3/13F8/N2	GATCATAGTATCCGCCGGAA (G/C) CCTAGGGAGCCCACAGCTCC
79	198	75	85/14B5/86	TTCGGCGGGTCGATCCGGGC [A/G]GAAGACATTGTCAGGTGANN
80	231	75	N1/14C2/N2*	GCACCAACATTGTAAACCTA [T/G] AGCTTCTTCCTCAGCCACCT
81	0	83	85/1402/86-1*	IGCTGCCACATAGTGAACCTA [T/A] AGCTTCTTCCTCAGCCACCT
82	33	83	N2/14C2/85-2*	GCACCAACATTGTGAACCTA (G/A) AGCTTCTTCCTCAGCCACCT
83	66	83	85/14C2/86-2	AGTACATAGCTATTGACTAA[C/G]TTAAGTTCCTTGTATTGTTG
84	99	83	N2/14C2/85-1	CCTCTATCCGCCATGGTTGC [A/T] CCAACATTGTGAACCTAGAG
85	132	83	85/14E2/86-2	TTGACCCTCGGCAAGCCACC[G/T]GTCAAGCCATGCTGCAGCCT
86	165	83	85/14E2/86-1	AGGCTGCCCTCTCCCAATTC (A/C) AAAGCCAACTCCTAAACCAA
87	198	83	185/14E8/86	AAACATGGAAAGGCCTGATA[/G]TCACCGTCAAGCTCACCGTC
88	231	83	85/14E12/86	CAACCTGAAAAATTGTTTTA (C/A) CAACGGCCCGGCTTTCTCCA
89	0	91	L1/14H10/86	AAGGCCAACAACGACATTAC (T/C) TCCATCGTTAGCAACGGAGG
90	33	91	85/14H10/86	TCACCGGCTTGAAGTCTTCC[G/T]CTGCATTCCCAGTCACCCGC

91	66	91	85/15A6/86	ACTCAGCTTTCTTATGCCTC [G/]ACTTGCGACACGAATCCA
92	99	91	85/15C4/86	TGCGGCTAACATCTCTGGTG (G/T) TCACCTTAACCCAGCCGTAN
93	132	91	85/15E5/86-1	CGAGGATCACTTCTCTGT (G/T) CAAGAAGAAGTTCGGCAAGG
94	165	91	N1/15E5/N2-1	CTGTtCAAGAAGAAGTTCGG [C/T] AAGGTCTACGCTTCCCGCGA
95	198	91	N1/15E5/N2-2	CCCTCTGCTCGTCACGGCGT T/A ACGCAGTTCTCGGATCTGAC
96	231	91	86/15E5/N2	CCCGCGAGGAGCACGACTAC [A/T]GATTCTCCGTTTTCAAATCC
97	0	99	L1/15E9/86*	TCCACTCGCCGGGAAGAAC (A/T)CGACAAACCGTTGTCTACTT
98	33	99	N2/15E9/L1	ATGGCTCGCGACGGGTCTCC (G/T)GTAAACCTCGGAGAGCAGAT
99	66	99	N2/15E9/86	GCCGACTCTCGAAGCTTCTT(A/)ACTCCACTCGCCGGGAAGAA
100	99	99	85/15E9/86-1	GAATCTAGGAGAGCAGATCT [T/G]CCTCTCTATCTTCAATGTTC
101	132	99	85/15E9/86-2*	TCCACTCGCCGGGAAGAAC [C/T]CGACAAACCGTTGTCTACAT
102	165	99	N1/15E9/N2-1	GTCATGAAGATATTCACTAC (A/G)CCGACTCTCGAAGCTTCTTA
103	198	99	85/15F1/86	GCAGGTAAAATTCTACAGAC (C/A)TTCCCTTTTCATTGTAGTTA
104	231	99	85/15F5/86	TCTCCTCCGCCGCCAAGAA (G/A)AAATCGACAGCGGCGCGTCT
105	0	107	85/15F10/86	GTGCCCTAAAGATACCCTCA (A/G)GCTTGGTGTCTGCGCTAATG
106	33	107	N2/15G1/L3	TTCTTCCCACAGGTGAAACT (T/C)GCTAACTTCCTTCCAAAGTA
107	66	107	IN1/15H7/N2	TATGTATCAGGACAATGTGT (GA/TT) GTGACTGTGGTTGCATCCAT
108	99	107	N1/16A1/N2-1	IGCTAAGCTACGCAACTGCCA (C/T) CAATCAGGGCAAGCTAAAGG
109	132	107	85/16A5/86	TATACACTCTTTAAAAGCGT [G/C]TGTGTGTACCCATCTCTCTT
110	165	107	N1/16B6/N2	ATGGCTGCGTATTGGCTGTC [C/T] AAGGCTGGATCTTGGTCCCA
111	198	107	85/16B6/N1	GGATCCATCTCAACTATGGT (A/C)GTATTATCGTTGAGGCTAGG
112	231	107	185/16B7/86	GTATGTGATTCGGAAGAGAA [T/]CAAACTAAGTGCCGAGAAAG
113	0	115	N1/16D6/N2	GCTAAGGTAGTTGGAGGAGC [CAA/GTG] CCACAGCCACGCGACTAAGG
114	33	115	85/16D10/86	CTCAACGTAGCAAGTAATAA [T/G]ATACTGTCTATTTATGGTTA
115	66	115	N1/16E9/N2	AGACTTTCCCCATTCTCTTC[T/A]CCATCCACCGTCGAAACCCA
116	99	115	185/16H3/86-1	ACTTCGAAACTGTAAACCTA (A/T)ACTTTAAGAGTTTAGAGCTA
117	132		B5/16H3/86-2	CACCATCGGAGAAAGAGGTA [C/T] TTCGAAACTGTAAACCTAAA
118	165		185/17A5/86	CTAAGGCGTCTCCTGAAGAA [A/G]TACAGAGAGTCGAAGAAGAT
119	198		185/1707/86	CCGCGGACGACGCTTTCTTC[C/A]TCTGCTCCACCGCGAGCGCC
120	231		185/17F7/86	GAGGAGTAGTCTCCATGGCC G/)AAGAAGAGCGTCGGAGACCTG
121	0		185/17G12/86	GAAGTTAGGGCTTCTAAGAT C/T AAGTTCGGCAAGGCTTTAAC
122	33		185/18A2/86	TCAAAACTAATATTTCTTTT[G/C]TTGATTGGTAATAAACAGGT
123	66		185/18A11/86	TTCCAGTGAAAAGGCATTGT T/G CTCCAAAATCTCGCTCTGCG
124	99		185/18F5/86	AAGCAGCTCTGACTTGAATG[C/A]GAGAGGTTAATCAGACTGTG
125	132		185/18H10/86-3	TAGATTGAAGCAATCAAGAA [G/A]ATCTCAGACTTCATCACCCA
126	165		185/19B3/86	GCATCCAACTCCAAGGATGA /C CCTGCCAAGGTGCTGCTAACT
127	198		B5/19C8/86	GAGCTCAGGGATGGTGGATC [A/T]GACTACCTTGGAAAGGGTCT
128	231		N1/19F4/N2	TGGGGTTAGTCGAAATAGGT (A/T)AAATGCTTTGAGTATGTGTA
129	0		N1/19H1/N2	TACGCGCAGCACGGACTTGC [G/A]ACGCAAGCAATCGAGCTTTT
130	33		85/20B4/86-1	GAAGCCCATGGTACGGAGCG [G/A] GAGAGAGTCAAGTACTTGGG
131	66		N1/20B12/N2	AACGGGTCACTGCTAAATCA (T/A) AAGGATCACAAGGCTGGGAC
132	99		85/20C12/86	CTAGCCTACTTTGGGAAAAG[/T]TTCGTTATTGTTTTGTGTCG
133	132	131		GACTTCAAGGACTTCGCCGG [A/C]AAATGCTCCGACGCTGTCAA
134	165		85/20D3/86-2	GAGGAGGCTACATGCAGCT[G/A]AAGAGGCTGAGGGGGCTAAA
135	198	131	85/20D6/86-4	GATGTTCAACCTATGAAGAA [G/C]AAACACCGAGGACCAACGAG
136	231	131	85/20D6/86-5	CCATTAGTGAGGGAGCATGT T / A CCTGTCACATTTGATGATTG

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137	0	139	85/20D6/86-8	AAACACATCGCCAAAGATCC[CG/AA]ACACTCGAGAAAGAGTGGAG
138	33	139	N1/20D8/N2	CTCATAGGCGATCTGGAGTA (T/G) GCAAATCGAATCTCCTCTCC
139	66	139	N1/20E1/N2	TGCACGCCTCACTTGTTCCT T / A CCAATCTGACATCAAGGATT
140	99	139	N1/20F1/N2-1	NGTGTTTTTGAGGTGAAAGC (A/T) ACAAATGGAGATACCTTTTT
141	132	139	N1/BoC-a2/N3-2	CCCGAGCCATTAGGACAAGA (T/C) GACTTGCCGTTTGACCAAAC
142	165	139	N1/BOC-A2/N3-1	CCCATCTCATCCTTTCTTGA [A / G] CCGTTGAATCAAGCTCCTGG
143	198	139	N1/BoC-a2/N3-3	TACATTCTCATTGGTTGGTT(C/A)TTGGGAAATAAAGTACCAAC
144	231	139	86/SC3/L2	GCACGCGCTAGAGTTGTTGC [C/A] AGAAGGAATGAACAATCTGA
145	0	147	N3/SC3/N4-1	CTTGAGACCTATAGTCCTGT (A/T) GTTCGGTCCGCCACAGTTCG
146	33	147	N3/SC3/N5-1	CACAGTTCGTACAGTTCTTC [A/C] CATTGCCACTGTTATGCACT
147	66	147	N1/SC3/N3-1	GAAGGCGTCCACTATCTTGA [A/G] ACCTATAGTCCTGTTGTTCG
148	99	147	86/SC3/N4-1	TCCCGGAAATCTTGCTGAAA (A/C)CGTTTACCTGCGACAACCAG
149	132	147	L3/B11/N5-1	ATGTCTTCAAAGTGCTCTGT[C/T]GCAACGCACGTCCGAACAAG
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	OLYMO	RPHIC	MARKERS	
			MARKER NAME	ISEQUENCE
150			s71/g2/g6-1	ATAATACTTGATATGCCATT[G/T]TGTCCTCTTATTTTTAACAT
151	19B		S67/G1/G3-3	GTACCTGCCGCCGCTGTCGA [CG/AC]GGACGACCTGCTGAAGCAGG
152	231		s67/g1/g5-1	ATGGCCTCGTCGGCCACTGC [A/C]GTCGCTCCGTTCCATGGGCT
253	0		S66/G1/G3-1	AATGTAATGGTACTCCGCGC T/C ATGGCTCTGGTACTTAGGAA
154	33		s65/g3/g6-2	ACCACTGACGTAGCACCTCC [G/T] ACTTCTCGTTGTAAAACCCC
155	66		S65/G3/G3-1	GGAGGTTCGCCTCATGTTAT (C/T)GTTGACGAGCCACATCCACT
156	99		S64/G1/g3-1	CTGGTTGAAATGTGTTGAAG[C/A]TACTAGTGATGAACTGCTTG
157	132		S63/g3/g6-1	TCTGTGATTGGAGTCTGCTC [G/A] CGTGTCAGCTCTGGATGTGA
158	165		S63/G4/G6-1	TACTGAGAGAATGCAACATC [C/G]AGCATTCTGTGATTGGAGTC
159	198		S57/G1/G2-4	GAAGCCAAATCCTATTATTT [T/C]CTGCCTCTAGGGTCTGAATG
160	231		S56/G4/G6-2	GCCTTATCATCCTCTAGGTA[T/A]TGGAGACGAGTGACCAGTCT
161	0		S56/G4/G6-1	GTACACTGTTACAATCACAC(T/G)TAGTGAAGCGCAACACAGAT
162	33		S53/G5/G6-5	GAGCGAGATCGATCCTGTTG[T/C]CATCCATCACTGCCATAGGA
163	66		S53/G4/G6-1	TAGTCATAGCAACAGCATGC (G/A) TCGTGATGTAGCGTTCACCC
164	99		S53/G1/G6-2	GAACAGAGTCCGCAATAGTT[T/C]ATCCTAATGCTACTTCGAGC
165	132		S53/G3/G5-1	CCCACGGCGGAGATGGTGG[T/]TAGAAGCGGAACCACCGAGC
166	165		549/G4/G5-1	CGGTGACCCGATGATTATGG[T/C]GGCGGCCACACCTGCAATGA
167	198			
	231		S49/G5/G1-2	CAAGCAAGCAAGCTGTCTGT (C/T) CGTATGTGTCTGGCATGTTA
168	231		S49/G2/G6-1	TGCTGCTGCACTTGCTCATC [G/C]TTACTGATTTGCTGAAATGT
169			549/G6/G1-3	TTTGCGCGCAATAAATCAGA (A/G)AGCTGATCTGAATCTGACCC
170	33		S49/G3/G6-1	CCACCCCGTTGCAGTGCTGT[/T]GCTGCTGCACTTGCTCATCG
171				GATTCAGAAACAGTGGCGGC (A/G) GATGTAGCATCAACACGCCC
172	99			CTCCTCGTGGTAGTGACGAT [G/C]ATTGCATCGGTGCCACAGGC
173	132			GGAATACTCGATAGGCTCCC[A/G]CTGTGGGTAACAGTATTCCT
174				TTCGGTGTCACTGACCTGTA (G/A) CATCAGCAGTAGCAGCGCCC
175				TCCGCGGAAACAACATCCGA [G/T]TTCTTGAGGATAACCCAGCT
176	231			ATGAGTATATTCAAGTCATA [T/C] TGTGAACTAGAATGTTATTT
177	0		S44/G5/G6-1	GCTGCGTCAATCATCACTTC (T/A) CCCACAGGCGTCAAGTACAG
178	33		S44/G1/G6-1	GCGTCAAGTACAGATACGCA (A/G)CACGCCTCAGCTTCGCCTTG
179	66	179	S43/G2/G6-2	CTTGATTGCATTGCAGCTAC [A/G]AGAAGCCCGTGGAAGGCCGG

132 179 843/G2/G3-1 CRGGTGGTGGTGCTTCTTCTG [ANAC/] TGAAACTGAACTGACTGCA 182 165 179 843/G1/G6-1 TGAATGGTAACCCACACTG [C/T] TGCGGTCGTTCTTGTGC 183 189 179 843/G2/G4-1 TGCAGTGGAACCTCCC [C/G] GAGAGGTTAACCCCACACTGTCGCT 185 0 187 843/G2/G5-1 TGCGGGAGGGTGAACCTCCC [C/G] GAGAGGTTCAACTGCGCTAGG 186 31 187 842/G3/G5-2 TGCGCGTAGGGTCCTCACCGT [G/C] GAGAGGTCAACTGCGCGAAGA 187 862/G2/G3-1 GCCGACAGGGTCCTCACCGT [G/C] GAGAGGTCCACTGAGAA 187 862 187 842/G2/G3-1 CTGCCTTCAGGAGGGCTCT [T/G] TGCGCTCGCGTCCATGAAA 188 99 187 842/G2/G7-1 GTTCTTGCACTCGATTGAGGGT [C/G] GAAGCAGGGCAGGGCGTGTC 188 99 187 842/G2/G7-1 GTTCTTGCACTCGATTGAGGGT [G/A] GGTCGGAAGCAGGGCAGGGCAGGC 190 165 187 841/G3/G1-1 CGCCAGATGACACCTCAC [TA/AGT] ATCCGGAAGAGAACCTAC 191 198 187 840/G2/G5-1 GATAGCCACATAGAACCTAC [TA/AGT] ATCCGGCAAGAGAACCCAC 193 195 840/G1/G4-2 GAGACCACCTCCGGA[G/A] ATACCGACCAACCCCCCAC 194 33 195 840/G1/G4-2 GAGACTAACAACCCTAC [TA/AGT] ATCCGCCAGGAACCCCCCCAC 194 33 195 838/G5/G7-2 ATAATGATGAACCCATC [G/A] CCGAAGAGATCCCCCCAC 195 195 195 195/G6/G2-1 GGTCAACAACCCATCCCGGAACCCACCCCCAC 197 132 195 197/G6/G2-1 GGTCAACAACCCATCCCGGAACCCACCCCCACCCCCACCCCCCACCCCCCACCCCCC					
182 165 179 S43/G1/G6-1 TGAATGGTAACCACAACTG [C/T] TCGCGTCCTGCTGGTTCATT 183 198 179 S43/G2/G5-1 TGCACGGGGGGTGAACCAACTGTCGCTT 185 0 187 S43/G2/G5-1 TACCCGGGGGGTGAACCACACTGTCGCTT 185 0 187 S43/G1/G5-1 GCCGACAGGGTCTACCCGT [G/C] AGCCCACACGATCATCGCCGAACACTGTTCGC 186 33 187 S42/G5/G7-2 TGCCGGACAGGGCGTC [T/G] TGCACCTGGGCTCCATCAAAACCGAGCGGAGACACCACACCACACCACACCACACCAC	180	99			GCTGGTTCATTATCTGACCT[G/T]GATTGCATTGCAGCTACAAG
198	181	132			
184 231 179 543/G2/G5-1 TACCCGGAGCTGAACCTCCC[C]GGAGAGATTCAAGTCGCTT 185 0 187 543/G3/G5-1 GCCGACAGGGTCCTCACCGT[G/C]AGCCCCTACAGCGAAGAA 187 543/G3/G5-2 TGCTGTCTCAGGGCGGCTGTC/GTAGACCAGAGAAA 187 542/G3/G3-1 CTGCACTCGGATTGAGGGGGCGTGTC/GTGAGACCAGGGCACCGCGTGTC 188 99 187 542/G3/G7-18 GTTCTCTGCACTCGATTGA[G/A]GGTCGGAAGCAGGGCACCGC 189 187 184 187 542/G3/G7-18 GTTCTCTGCACTCGATTGA[G/A]GGTCGGAAGCAGGGCACCGC 189 187 187 541/G3/G3-1 GTTCTCTGCACTCGATTGA[G/A]GGTCGGAAGCAGGGCACCGC 189 187 184 187 541/G3/G3-1 GTTCTCTGCACTCGATTGA[G/A]GGTCGGAAGCAGGGCACCGC 189 187 540/G3/G3-1 GGCCGAGATGACCACTGCA[TA/AGT]ATCCTGCAAGCAAGGAGATACCCT 189 187 540/G3/G4-2 AAGTACCACTTGAGACCTTGA[G/A]ATACGGCCAAGAGAATCCCT 189 187 540/G3/G4-2 GGAGCTTAAGGATCCCTCCGGA[G/A]ATACGCCAAGACACCTTG 189 185 538/G5/G7-2 AAGTACGACACCTCCGGA[G/A]ATACGGCCAACCACCTTG 189 185 538/G5/G7-2 ATAATGATACCACTCCGGA[G/A]ATACGCCAACCACCTTG 189 185 538/G5/G7-2 ATAATGATACCTCTCGGA[G/A]ATACGCCAACCACCTTG 186 185 538/G5/G7-2 ATAATGATACTCTTGAG[G/T]AGATTCAAAGACAACCTTG 186 185 537/G3/G3-2 GGCAATAAGGATACTCTGG[G/A]ATACGACCACCGGACGACGACGACGACGACGACGACGACGAC	182	165	179	IS43/G1/G6-1	
185	183	198	179	IS43/G2/G4-1	
186 33 187 \$42/G5/G7-2 TCCCTGTCAGGGAGGGCGTC T/G TGCAGCTCGATTGAAA 187 66 187 \$42/G2/G7-1 GTCCACTCGATTGAGGATC C/G GAAGCAGGGCAGGCGTGTC 188 99 187 \$42/G2/G7-1 GTCCTGCACTCGATTGAGGATC C/G GAAGCAGGGCAGGCCGTGTC 189 132 187 \$42/G2/G7-1 GTTCTCTGCACTCGATTGA G/A GGTCGAGAGCAGGGCAGGCC 190 165 187 \$41/G3/G1-1 GCCCAGATGAACCACTGA G/A GGTCGAGAGCAGGGCAGGCC 191 198 187 \$41/G3/G1-1 GCCCAGATGAACCACTCGA G/A GGTCGAGAGCAGGAGAGCAGGCC 191 198 187 \$41/G3/G1-1 GAAGCCACATAGAACCATTG G/T) CCTCCAGCTAACAACAGAG 192 231 187 \$40/G4/G2-1 AAGTCACACACAGAG G/A AATTCCCTGCAGAGAATCCCGC 193 0 195 \$40/G4/G2-2 GAGGCTTAACGATACCACTG GA/CACAACAGAG 194 33 195 \$38/G5/G7-2 AATACATGGAACCATACT GA/CCGAAGATGACACACACTT 195 66 195 \$38/G5/G7-2 ATACATGGAACCATACT GA/CCGAAGATGAACACACACTT 196 99 195 \$37/G2/G3-2 TCCAGGGGGCACCACACTT GA/CCGAAGATGAACACACACTT 197 132 195 \$37/G2/G3-2 TCCAGGGGGCACCGAAGGG AATCCAAGACATGAGACACTTCC 198 165 195 \$37/G2/G3-2 TCCAGGGGGCACCGAAGGG AATCCAAGACACACACTTCC 199 198 195 \$37/G1/G6-3 AATCCATCACCCGAACAGGCG/AACACACACACTTCC 199 198 195 \$37/G1/G6-2 AATTCAGCTCAAATCATAGGI ATCCAAGACACACACACACACACACACACACACACACACA	184	231	179	IS43/G2/G5-1	
187 66 187 \$42/G2/G3-1 CTGCACTCCGATTGAGGGTC C/G GAAGCAGGGCAGCGCCTGTC 188 99 187 \$42/G2/G7-1b GTTCTTCTGCACTCCGATTGA G/A GGTTCGAAGCAGGGCAGCGC 189 132 187 \$42/G2/G7-1b GTTCTTCTGCACTCCGATTGA G/A GGTTCGAAGCAGGGCAGCGC 180 145 187 \$41/G3/G1-1 CGCCGAATGAACACCTCCGATTGA G/A GGTCGAAGCAGGGCAGCGC 191 198 187 \$41/G3/G1-1 CGCCGAATGAACCACTCTCA G/A GGTCGAAGCAAGAACACAGAG 192 231 187 \$40/G1/G2-1 AAGTACACACTCCGGA G/A ATACGGCCAAACAACAGAG 193 0 195 \$40/G1/G4-2 GGAGCTTAACGACCTCCGGA G/A ATACGGCCAAAGAACACAGAG 194 33 195 \$38/G5/G7-2 ATAATGATGGACCACTCCGGA G/A ATACGGCCAAAGAACCATT G/T CGCAAGAAGAACCATT 195 66 195 \$38/G5/G7-2 ATAATGATGGAACCATT G/A CCGAAGATGAACCACCTTG 196 99 195 \$37/G6/G2-1 GGTCAATAAGATACACA G/A AAAACCTTCGGTGAAGCACTCGGA 197 132 195 \$37/G6/G2-1 GGTCAATAAGATACACA G/A AAAACCTTCGGTGAACCACCCGAGGCA 198 165 195 \$37/G6/G5-1 GGTCAATAAGATACACA G/A AAAACCTCACGAACA 199 198 195 \$37/G1/G6-2 AATACGATGACCACACAGAGG AAAACGACCACCACAGAGG 199 198 195 \$37/G1/G6-2 AATACGATGACCACACCAGACAG G/T CACTGGCACACCACACAGAACA 200 231 195 \$37/G1/G6-2 AATACGATCACACAACACCACACAGACAG G/T CACTGGCACACCCACACACACACACACACACACACACACA	185	0	187	IS43/G1/G5-1	GCCGACAGGGTCCTCACCGT G/C AGCCCCTACTACGCCGAAGA
188 99 187 \$42/\$\text{G2}\$\text{G7-1a} \text{GTTCTGCACTCCGATTGA} \text{G/A} \text{GGAGGGCAGGCCC} \\ 189 132 187 \$42/\$\text{G2}\$\text{G7-1b} \text{GTTCTGCACTCCGATTGA} \text{G/A} \text{GGTCAGGGCAGGCCAGGCCC} \\ 190 165 187 \$41/\$\text{G3}\$\text{G1} \text{GGCCAGAGGACCACTCAGTCAGTCAGTCAGTACAGGGGAAGGAGATATCCCT} \\ 191 198 187 \$41/\$\text{G2}\$\text{G5}\$\text{-1} \text{GAGGACCACTAGAACCAGTAT} \text{G/T} \text{CCGGAGAACACCAGGG} \\ 192 231 187 \$40/\$\text{G4}\$\text{G2}\$\text{-1} \text{AAGGACCACTAGAACCATGGA} \text{G/A} ATACCCCGCACTAGAACCACGGGCACACCAGAGATCCGGCACCACCAGAGATCCGCACCACCACCACCACCACCACCACCACCACCACCACC	186	33	187	S42/G5/G7-2	TCGCTGTCAGGGAGGGCGTC T/G TGCAGCTCGGCTCCATGAAA
189 132 187 542/G2/G7-1b GTTCTCTGCACTCCGATTGA[G/A]GGTCGGAAGCAGGGCAGCGCC 190 165 187 541/G3/G5-1 GGCCGAATGAACCAGCTCAC(TA/AGT)ATCCTGCAAGCAACCAGAGAGATATCCCT 191 198 187 541/G2/G5-1 GATAGCCACATAGAACCAGCTCAC(TA/AGT)ATCCTGCAACCAACAACGAG 192 231 187 540/G4/G2-1 AAATCACTCCGGA GATAGCCACATAGAACCTCTGGA G/A)ATACCGCACAACAACCGAG 193 0 195 540/G4/G2-1 AAATCACACCTCCGGA GATAGCACATGGACTGGA GATAGCACACCTGGA GATAGCACACCACCTGGA GATAGCACACCACCACCCTG GATAGCACACCACCACCCCGGCACCCCGAGAGGC AAACCCACCACCACCACCCACCCCGAGAGCC GATAGCACACCACCACCACCACCACCCACCCCGACCCCCACCCCCACACCCCCACCCCCACCCCCACCCCCACCCC	187	66	187	IS42/G2/G3-1	CTGCACTCCGATTGAGGGTC C/G GAAGCAGGGCAGCGCGTGTG
190	188	99	187	S42/G2/G7-la	GTTCTCTGCACTCCGATTGA [G/A]GGTCcGAAGCAGGGCAGCGC
191 198 187 S41/G2/G5-1 GATAGCCACATAGAACCTTG[G/T]CCTCCAGCTAACAACACGAG 192 231 187 S40/G4/G2-1 AAGTACATGCACCTCCGGA [G/A)ATACGGCCAAGAATCCGGCA 193 0 195 S40/G1/G4-2 GGAGCTTAACGATC[GA/C]AAATTCCCTCCCCGGCACACATTGCATCCTTCCCGGA [G/A)ATACGCCCAGGACATGCAATCGCACATTGCATCCTTCCTCCCCCGGCACACATTGCATCCTTCCT	189	132	187	S42/G2/G7-1b	GTTCTCTGCACTCCGATTGA [G/A] GGTCGGAAGCAGGGCAGCGC
192 231 187 S40/G4/G2-1 AAGTACATGACACCTCCGGA [G/A] ATACGGCCAAGAATCCCGCA 193 0 195 S40/G1/G4-2 GGACCTTAACGATACCAATC [GA/CG] AAATTCCCTCGCGGGCAC 194 33 195 S38/G5/G7-2 ATAATGATGGAACCATGACT [GA/CG] AAATTCACCACCTTG 195 66 195 S38/G5/G7-1 GAGATATATGATGTACTTCTAG [G/T] AGATTCAAAGCAACCTTG 196 99 195 S37/G6/G2-1 GGTCAATAAGATAACTACAC [C/A]AAACTCTGCGTACAGTCTG 197 132 195 S37/G2/G3-2 TGCAGGCGGCGCGCAAGCGT [C/A]AAACTCTGCGTACAGTCTG 198 195 S37/G2/G3-2 TGCAGGCGGCGCAAGCGG [C/A]ATCGAGGCCTCACCGGTT 198 195 S37/G1/G6-3 CAATACGATCAGCCAGACCG [C/A]ATCCGGGGTGTCCCAGAGG 199 198 195 S37/G1/G6-3 CAATACGATCAGCCAGACAC [C/A]CACTGAGCAACATCAGCAAA 200 231 195 S37/G1/G6-2 AATTCAGCTCAGAACCCAGACAC [C/A]CACTGACAATCACACCAA 201 0 203 S37/G2/G4-4 GTCTCCCGGATCCAGACAC [C/A]CACTACACCAAA 202 33 203 S36/G3/G6 TCACTTTTATGTGGTGTTTA [C/G]CATAAGGATAACTACACCAA 203 66 203 S35/G2/G3-2 GCTCGAACTTCATGTTTA [C/G]CATGATGTGCTACATCGCTTTC 204 99 203 S35/G3/G6-1 TCTGAGATCATTCATTTAT [A/G]CATGATGTGTATTATCTTTC 204 99 203 S35/G3/G6-1 GCTGGAACTTCATTTATTCATTTA [C/A)ATGTCAGATAAAGAAAACTC 205 132 203 S34/G3/G5-2 GGTTGGACCATATCATGAT [C/T] GGAATGCCGCCTCAAAATGC 206 207 198 203 S34/G3/G5-2 GGATTGCACCATATCATGAT [C/T] GTATGATCTTTACTCTCC 207 198 203 S34/G3/G5-2 GGATTGCACCATATCATGAT [C/T] GTATGATCTTTCCTTTTTTTTTTTTTTTTTTTTTTTTTT	190	165	187	S41/G3/G1-1	CGCCAGATGAACCAGCTCAC [TA/AGT] ATCCTGCAAGAGATATCCCT
193 0 195 \$40/G1/G4-2 GGAGCTTAACGATACCATC [GA/CG]AAATTCCCTCGCGGGCAC 194 33 195 \$538/G5/G7-2 ATAATGATACCATG [GA/CG]AAATTCCCTCGCGGGCAC 195 \$538/G5/G7-2 GAGATATATGGTACTTCTAG [G/T]AGATTCAAGACCAAGCTTG 195 66 195 \$537/G6/G2-1 GGTCAATAAGATAACTACAC [C/A]AAACTTGCGTACAGTCTCC 197 132 195 \$537/G6/G2-2 TGCAGGCGGCAGCGGAAGCG [A]AATCGAGCCTGACCGGTCT 198 165 195 \$537/G4/G5-2 ATCGATGGACAAGCGGGAAGCG [A]AATCGAGCACAGCCTGACCGGTGT 198 195 \$537/G1/G6-3 CAATACGATCAGCCAAGACG [C/T]CACTGGCAGACATCCAGCAAA 195 \$537/G1/G6-2 AATTCAGCTCAAATCATAGG [T/G]CAATAAGATAACTACCAAA 195 \$537/G1/G6-2 AATTCAGCTCAAATCATAGG [T/G]CAATAAGATAACTACCAAA 195 \$537/G1/G6-2 AATTCAGCTCAAATCATAGG [T/G]CAATAAGATAACTACCCAAA 195 \$537/G1/G6-2 AATTCAGCTCAAATCATAGG [T/G]CAATAAGATAACTACCCAAA 196 196 195	191	198	187	S41/G2/G5-1	GATAGCCACATAGAACCTAT [G/T] CCTCCAGCTAACAACACGAG
194 33 195 S38/G5/G7-2 ATAATGATGGAACCATGACT [G/A] CCGAAGATGAACCAACCTTG 195 66 195 S38/G5/G7-1 GAGATATATGGTACTTCTAG [G/T] AGATTCAAAGACATGGAACCA 196 99 195 S37/G5/G5-2 GGTCAATAAGATAACACA [C/A] AAACCTTGCGTACAGTTCC 197 132 195 S37/G2/G3-2 TGCAGGCGGAGCGGAGCGG [A] ATCGAAGGCCTGACCGTTCTC 198 165 195 S37/G1/G6-3 ATCGATGGACAAGGCAGGGT [A/G] ATCTCGGGGTGTGCCAGAGG 199 198 195 S37/G1/G6-3 CAATACGATCAGCCAACCCCAACACCACCAACCCCAACCCCAACCCCACCCACCCC	192	231	187	S40/G4/G2-1	AAGTACATGACACCTCCGGA [G/A] ATACGGCCAAGAATCCCGCA
195	193	0	195	S40/G1/G4-2	GGAGCTTAACGATACCAATC [GA/CG] AAATTCCCTCGCCGGGCAC
196 99 195 537/G6/G2-1 GGTCANTAGGTACACC C/A AAACTCTGCGTACAGTCTCC 197 132 195 537/G2/G3-2 TGCAGGCGGCAGCGGAGCG /A ATCGATGGCCAGAGG /A ATCGATGGCCAGAGGCGT /A ATCGATGGCCAGAGAGCG /A ATCGATGGCCAGAGAGCG /A ATCGATGGCCAGAGACAGCGCT /A ATCGATGACAACATCAGCAAA CONTACGACAACATCAGCAAA CONTACGACAACATCAGCAAA CONTACGATCAGCAGACAGCCAGACAGCCGT /A ATCGATGAGATACATCAGCCAA CONTACGATGAGCAGACATCAGCACACACCACCACACACCACCACACACA	194	33	195	IS38/G5/G7-2	ATAATGATGGAACCATGACT [G/A]CCGAAGATGAACCAACCTTG
197 132 195 537/G2/G3-2 TGCAGGCGGCAGCCGAAGCG[/A]ATCGAAGGCCTGACCGGTCT 198 165 195 537/G4/G5-1 ATCGATGGACAAGGGAGGGT[A/G]ATCTCGGGGTGTGCCAGAGG 199 198 195 537/G1/G6-2 CAATACGATCAGCAAACCTCAGCAACCTCAGCAACCTCAGCACATCCTCCGGATCCAGAATCCTCAGCAACCTCAGCACATCTCCGGATCCAGAATCCTCCGGATCCAGAATCCTCCGGATCCAGAATCCTCCGGATCCAGAATCCTCTTACACCCAACCTCTCCGGACATCCTCAGATCACCACAACCTCCAGAATCCTCTTAGTGGTTTTACCGGACATCCTCTTCCCGGACTCTAGATGGATG	195	66	195	538/G5/G7-1	GAGATATATGGTACTTCTAG [G/T] AGATTCAAAGACATGGAGCA
198	196	99	195	S37/G6/G2-1	GGTCAATAAGATAACTACAC [C/A]AAACTCTGCGTACAGTCTCG
198 195	197	132	195	S37/G2/G3-2	TGCAGGCGGCAGCCGAAGCG[/A]ATCGAAGGCCTGACCGGTCT
200 231 195 S37/G1/G6-2 AATTCAGCTCAAATCATAGG[T/G]CAATAAGATAACTACACCAA 201 0 203 S37/G2/G4-4 GTCTCCCGGATCCAGAATCC[C/T]ACCGCGTTACATCGCCTTTC 202 33 203 S36/G3/G6 TCACTTTTAGTGGTGTTTA[C/G]ACTAAGGATGCTGACATCT 203 66 203 S35/G2/G3-2 GGTCGAACTTCTAGTTGATT[A/G]CATGATTGCTATTACTGTTG 204 99 203 S35/G3/G6-1 TCTGAGATACATTCATTTAA[C/A]ATGCCAGATAAAGAAAAATCC 205 132 203 S35/G3/G6-1 TCTGAGATACATTCATTTAA[C/A]ATGCCAGCCCATAAAATGC 206 165 203 S34/G3/G5-2 GGTTTGCACCCATATCATGAT[C/T]AGTTGATCTTTTATTCTCC 207 198 203 S34/G3/G5-2 GATTTTGTAGGTTGATGCAT[C/T]GTTTGATCTTTTATTCTCC 208 231 203 S33/G4R/G6-1 GGCAGACAACAGACAGACCATCA[A/C]GCATCGTTGCATTTACTCTCA 209 0 211 S33/G4/G2-1 GGCAGACAACAGACAGACAGACCAGATCA[A/C]GCATCGTTGCATTTACTCTCA 210 33 211 S33/G5/G6-2 TCAAAGTGGTGCAATCGCAA[T/C]CCACTTGGGCTTGCCGTGGT 211 66 211 S33/G2R/G3-1 ACGCATGCTTGCATTTACTC[C/T]CAGTCAAAACTCAA 212 99 211 S32/G3/G5-2 AATGAACACACACATCATCCTCTAAAAACTCAA 213 132 211 S32/G3/G5-2 AATGAATCACACATTCTTCCGAA[A/G]CGTACCACAAAACTCGA 214 165 211 S31/G3/G5-1 CAAAGTCACCATTCTTCCGAA[A/G]CGTAACACCACATTCTTCCGA 215 198 211 S31/G3/G6-1 TGTATCAGCATTGTGTATACA[A/G]CGAAAGTTGAATCCCCACA 216 231 211 S29/G4/G5-1 TGTATCAGCATGTTGAGAT[G/A]TCGCACTGGACTCAGTGCT 217 0 219 S29/G1/G5-1 CCATTGCTTGTTATCACATACCACACTTCTTCCACA 219 66 219 S28/G3/G6-1 GGAATGTTGCTTATATTAC[T/C]CGTAGGTGACACAGGCTTCACGACCATTCTTCCACACACA	198	165	195	S37/G4/G5-1	ATCGATGGACAAGGGAGGGT A / G ATCTCGGGGTGTGCCAGAGG
201 0 203 S37/G2/G4-4 GTCTCCCGGATCCAGAATCC [C/T]ACCGCGTTACATCGCCTTTC 202 33 203 S36/G3/G6 TCACTTTTTAGTGGTGTTTA [C/G]ACTAAGGATGCTGACATTCT 203 66 203 S35/G2/G3-2 GCTCGAACTTCTAGTTGATT [A/G]CATGATTGCTATTACTGTTG 204 99 203 S35/G3/G6-1 TCTGAGATACATTCATTTAA [C/A)ATGTCAGATAAAGAAAACTC 205 132 203 S35/G4/G5-1 GGTTCGCACCATATCATGAT [C/T]GGAATGCCGCCTCAAAATGG 206 165 203 S34/G3/G5-2 IGATTTTGTAGGTTGATGCAT [C/T]GGAATGCCGCCTCAAAATGG 207 198 203 S34/G2/G5-1 TGTAGGACTTGGAGACCATG [A/G]TAATTTACACTCCCTGT 208 231 203 S33/G4R/G6* IGGCAGACAACAGACGATCA [A/C]CATGCTTGCATTTACTCTCC 209 0 211 S33/G4R/G6* IGGCAGACAACAGACAGATCA [A/C]CACTGCTTGCATTTACTCTCA 210 33 211 S33/G5/G6-2 ITCAAAGTGGTGCAATTACTC [C/T]CAGTCATGCATTTACTCTCA 211 66 211 S33/G2R/G3-1 ACGCATGCTGCATTTACTC [C/T]CAGTCAAAACTCGA 212 99 211 S32/G3/G5-2 IACGAATGCAATTCCCGAA [A/C]CTGCTTGCCTTGCCGTGGT 213 132 211 S32/G3/G5-2 IACGAATGCAATA [C/T]AGAATCCACCATTCTCCGA 214 165 211 S31/G3/G5-1 ICAATGTCTTGTTTCTGAATA [C/T]AGAATCCACCATTCTTCCGA 215 198 211 S31/G3/G4-1 ITCTATCAGCATTTACTC [C/T]AGAATCACCATTCTTCCGA 216 231 211 S29/G4/G5-1 ICAATGTCTTGTTACTCTGGATGGAATGCCACAC 217 0 219 S29/G1/G5-1 ICCATGCGCTAGTCTGGATGGAATGCCTGAATGCCTGAATGCCTGAATGCCTGAATGCCTGAATGCCTG 218 33 219 S28/G2/G6-1 IGAATGTTGCTTTATACACGCC [T/T]CAGATGTGTGGGATTCACACAAACTCCGAA 220 99 219 S28/G3/G4-2 IACGCATCTGTTATATACCTCCCAAAACTCTGAATGCCTG 221 132 219 S28/G3/G4-2 IACGCGGGGGCGCACTCGAGTGACACACACACACACACACA	199	198	195	S37/G1/G6-3	CAATACGATCAGCCAGACAG [C/T] CACTGTGCAACATCAGCAAA
202 33 203 536/G3/G6 TCACTTTTTAGTGGTGTTTA[C/G]ACTAAGGATGCTGACATTCT 203 66 203 S35/G2/G3-2 GCTCGAACTTCTAGTTGATT [A/G]CATGATTGCTATTACTGTTG 204 99 203 S35/G3/G6-1 TCTGAGATACATTCATTTAA[C/A]ATGTCAGATAAAGAAAACTC 205 132 203 S35/G4/G5-1 GGTTCGCACCATATCATGAT[C/T]GGAATGCCGCCTCAAAATGG 206 165 203 S34/G3/G5-2 GATTTTGTAGGTTGATGCT[C/T]GTATGACTTTCTTATCTCC 207 198 203 S34/G2/G5-1 TGTAGGACTTGGAGAGCTTG[A/G]TAATTTACACATGCCTCTGT 208 231 203 S33/G4R/G6-1 GGCAGACAACAGACAGATCA[A/C]CATGCTTGCATTTACTCTCA 209 0 211 S33/G4R/G2-1 GGCAGACAACAGACAGATCA[A/C]CATGCTTGCATTTACTCTCA 210 33 211 S33/G5/G6-2 TCAAAGTGGTGCAAT[T/C]CCACTTGGCGTTGCCTTGGTT 211 66 211 S33/G2R/G3-1 ACGCATCCTTGCATTTACTCTCCA 212 99 211 S32/G3/G5-3 GAATCCACCATTCTTCCGAA[A/G]CTGCTTCCTACAAAACTCGA 213 132 211 S32/G3/G5-2 ATGAATTGAAGCTCTGAATA[C/T]AGAATCCACCATTCTTCCGA 214 165 211 S31/G3/G5-1 CAATGCTTGTTCTTATCA[A/G]CGAAAGTTGAATCCCCACA 215 198 211 S31/G3/G4-1 TGTATCGGCTAGTTGATCA[A/G]CGAAAGTTGAATCCCCACA 216 231 211 S29/G4/G5-1 CAATGCTTGTTCGTTATCA[A/G]CGAAAGTTGATCCCCACA 217 0 219 S29/G1/G5-1 CCTTACACTATTTACAGGCC[C/T]AGGATGCTGGGTTCCTTCAG 218 33 219 S28/G2/G6-1 GAATGTTGTTGTTATATATCC[C/T]AGGATGTTGCGTTCTTCAG 219 66 219 S28/G3/G6-1 GAATGTTGTTTATATATATCT[T/C]CGTAGTGTACAAAGGGTTCA 220 99 219 S28/G3/G6-1 GAATGTTGTTTATATATATCTT/C]CGCAGAGCAGCGAAGCTTCA 221 132 219 S28/G3/G6-1 GAATGTTGTTCTTCAGAACCTTTTTTTCAGACCTTTATACACTGAAAACCTTT 222 165 219 S28/G3/G6-2 ITCCGTTGTTATCAGAGCC[T/T]TAAAATATTGTTTCAAACTTT 222 165 219 S28/G3/G6-2 ITCCGTTGTTATCACTGAACCTTTTTTTTTCAAACTTT 222 165 219 S28/G3/G6-2 ITCCGTTGTTGTTCTTCAGTTCTTAAAATATTGTTTCCAAGCCCAAAGCCTTTTAAAAATATTGTTTTCCAAGCCTTAAAAATATTGTTTCCAAGCCCAAAGCCAAAGACCAAAGAAAAAAAA	200	231	195	S37/G1/G6-2	ARTTCAGCTCAAATCATAGG[T/G]CAATAAGATAACTACACCAA
203 66 203 S35/G2/G3-2 GCTCGAACTTCTAGTTGATT [A/G]CATGATTACTGTTG 204 99 203 S35/G3/G6-1 TCTGAGATACATTCATTTAA [C/A)ATGTCAGATAAAGAAAACTC 205 132 203 S35/G4/G5-1 GGTTCGCACCATATCATGAT [C/T]GGAATGCCGCCTCAAAATGG 206 165 203 S34/G3/G5-2 IGATTTGTAGGTTGATGCAT [C/T]GGTAGCCTCTAAAATGG 207 198 203 S34/G2/G5-1 TGTAGGACTTGGAGAGACTC [A/G]TAATTTACACATGCCTCTGT 208 231 203 S33/G4R/G6- IGGCAGACAACAGACAGATCA [A/C] GCATGCTTGCATTTACTCTCA 209 0 211 S33/G4F/G2- IGGCAGACAACAGACAGATCA [A/C] GCATGCTTGCATTTACTCTCA 210 33 211 S33/G5/G6-2 TCAAAGTGGTGCAATTCATC [C/T]CAGTCAAAACTCAGTCCCGGAT 211 66 211 S33/G2R/G3-1 AGGCATGCTTGCATTTACTC [C/T]CAGTCAAAACTCAGACAGACAGACAGATCA [A/C] GCATGCTTGCCGTTGCT 212 99 211 S32/G3/G5-3 IGAATCCACCATTCTTCCGAA [A/G] CTGCTTCCTACAAAACTCGA 213 132 211 S32/G3/G5-2 ATGAATTGAAGCTCTGAATA [C/T]AGAATCCACCATTCTTCCGA 214 165 211 S31/G3/G5-1 ICAATGTCTTGTTATCA [A/G] CGAAAGTTGAACCCACA 215 198 211 S31/G3/G4-1 ITGTATCGGCTAGTTGTTACTA [A/G] CGAAAGTTGAATCCCCACA 216 231 211 S29/G4/G5-1 ICCAATGTCTTGAGAA [G/A]/TCGCACTTCAGTGCT 217 0 219 S29/G1/G5-1 ICCATACACTATTACTC [C/T]AGGATGTCTGAATGCCTG 218 33 219 S28/G2/G6-1 IGAATGTTGCTTATACTAC [T/C] CGTAGGTGACAAAGGGTTCA 220 99 219 S28/G3/G6-1 IGAATGTTGCTTATATTAC [T/C] CGTAGGTGACAAAGGGTTCA 220 99 219 S28/G3/G6-1 IGAATGTTGCTTATATTAC [T/C] CGTAGGTGACAAAGGGTTCA 221 132 219 S28/G3/G6-2 IGCGTAGCTTCCCGAACACTGTTTTTTTAACAAGCC [T/T] TCAACACTATTATTCAAACTTT 222 165 219 S28/G3/G6-2 ITCACTTTAACAGCC [T/T] TAACACTTTTATTCAAACTTT 222 165 219 S28/G5/G6-2 ICAAGAAGCCTCTTCAGTGTC [A/C] IGTCGTAGCTTCCTCAAGACC 223 198 219 S27/G3/G6-2 ITCCCTTGGTAGCCTTCAGTGTC [A/C] ITTAACACTTTTATCCAAGACC 224 231 211 219 S26/G5/G6-2 ICCAAGAAAGATTAATGCTTG [T/TT] TAAAAATATTGTTTCCAAGACC 224 231 219 S26/G5/G6-2 ICCAAGAAAGATTAATGCTTG [T/TT] TAAAAATATTGTTTCCAAGCCCAAGCC 224 231 29 S26/G5/G6-2 ICCAAGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	201	0	203	S37/G2/G4-4	GTCTCCCGGATCCAGAATCC (C/T) ACCGCGTTACATCGCCTTTC
204 99 203 \$35/G3/G6-1 TCTGAGATACATTCATTTAA(C/A)ATGTCAGATAAAGAAAACTC 205 132 203 \$35/G4/G5-1 GGTTCGCACCATATCATGAT(C/T)GGAATGCCGCCTCAAAATGG 206 165 203 \$34/G3/G5-2 GATTTTGTAGGTTGATGCAT(C/T)GTTTGATCTTCTTATCTCC 207 198 203 \$34/G2/G5-1 TGTAGGACTTGGAGAGCTTG(A/G)TAATTTACACATGCCTCTGT 208 231 203 \$33/G4R/G6-1 IGGCAGACAACAGACAGATCA(A/C)CATCCTTGCATTTACTCTCA 209 0 211 \$33/G4R/G6-2 IGGCAGACAACAGACAGATCGCAA(T/C)CCACTTGCGTTGCCGTGGT 210 33 211 \$33/G5/G6-2 TCAAAGTGGTGCAATCGCAA(T/C)CCACTTGCATTGCCGTGGCTTGCCGTGGT 211 66 211 \$33/G2R/G3-1 ACGCATGCTTGCATTTACTCTC/C/T)CAGACACCCACTTCTCCGAA 212 99 211 \$32/G3/G5-2 ATGAATTGAACCTCTGAATA(C/T)AGAACCCACTTCTCCGAA 213 132 211 \$31/G3/G5-1 CAATGTCTTGTTCTGTTATATCA(A/G)CGAAAAGTTTGAATCCCCACACTTCTCCACACTTCTCCACACTTCTCAC	202	33	203	536/G3/G6	TCACTTTTTAGTGGTGTTTA [C/G]ACTAAGGATGCTGACATTCT
205 132 203 S35/G4/G5-1 GGTTCGCACCATATCATGAT [C/T] GGATTGCCGCCTCAAAATGG 206 165 203 S34/G3/G5-2 GATTTGTAGGTTGATGCAT [C/T] GTTTGATCTTTCTTATCTCC 207 198 203 S34/G2/G5-1 TGTAGGACTTGGAGAGCTTG [A/G] TAATTTACACATGCCTCTGT 208 231 203 S33/G4R/G6 IGGCAGACAACAGACAGATCA [AG/CA] CATGCTTGCATTTACTCTCA 209 0 211 S33/G4T/G2 IGGCAGACAACAGACAGATCA [A/C] GCATGCTTGCATTTACTCTCA 210 33 211 S33/G5/G6-2 ITCAAAGTGGTGCAATCACAAA [T/C] CCACTTGGGCTTGCCGTGGT 211 66 211 S33/G2R/G3-1 ACGCATGCTTGCATTTACTC [C/T] CAGTCAAACTCAGACCGAA 212 99 211 S32/G3/G5-2 GAATCCACCATTCTTCCGAA [A/G] CTGCTTCCTACAAAACTCGA 213 132 211 S32/G3/G5-2 ATGAATTGAAGCTCTGAATA [C/T] AGAATCCACCATTCTTCCGA 214 165 211 S31/G3/G4-1 ICCATTCTTCTTCTTATCAA [A/G] CGAAAGTTTGAATCCCCACA 215 198 211 S31/G3/G4-1 ITGTATCGGCTAGTCTGGATG [G/A] TCGCACTGGCACTCAGTGCT 216 231 211 S29/G4/G5-1 ICCTTACACTATTAACAGGCC [C/T] GTGATCTACCTGAATGCCTG 217 0 219 S29/G1/G5-1 ICCTTACACTATTAACAGGCC [C/T] GTGATCTACCTGAATGCCTG 218 33 219 S28/G3/G6-1 IGCGTAGCTTCCTCAAGACC [T/T] TCCTGATGTGCGGACCAAGCCTCAA 220 99 219 S28/G3/G6-2 IAGGGGGTGGTGCGGACTGGA [T/G] CGCCCGAGCAGCCGAAGCC 221 132 219 S28/G3/G6-2 IAGGAGGCTGCTGCGGACTGGA [T/G] CGCCCGAGCAGCCGAACCC 223 198 219 S28/G3/G6-2 ICCAAGAAGCCTTTCAGTGTC [A/C] GTCGTAGCTTCCTCAAGACC 224 231 219 S26/G5/G6-2 ICCAAGAAGCCTTTCAGTGTC [A/C] GTCGTAGCTTCCTCAAGACC 224 231 219 S26/G5/G6-2 ICCAAGAAGACTTTCAGTGGC [T/T] TAAAATATTGTTTCCAGCCACACCCACCCCACCCCCACCCCCACCCCCACCCCCACCCC	203	66	203	535/G2/G3-2	GCTCGAACTTCTAGTTGATT [A/G] CATGATTGCTATTACTGTTG
206	204	99	203	S35/G3/G6-1	TCTGAGATACATTCATTTAA (C/A)ATGTCAGATAAAGAAAACTC
207 198 203 S34/G2/G5-1 TGTAGGACTTGGAGAGCTTG[A/G]TAATTTACACATGCCTCTGT 208 231 203 S33/G4R/G6* IGGCAGACAACAGACAGATCA[A/C]GCATGCTTGCATTTACTCTCA 209 0 211 S33/G4r/G2* IGGCAGACAACAGACAGATCA[A/C]GCATGCTTGCATTTACTCTCA 210 33 211 S33/G5/G6-2 TCAAAGTGGTGCAATTCCCAATTCCCACTTGGGCTTGCCGTGGT 211 66 211 S33/G2R/G3-1 ACGCATGCTTGCATTTACTC[C/T]CAGTCAAACTCAGACACAGACAGACAGATCA[A/C]GCACTTGGGCTTGCCGTGGT 212 99 211 S32/G3/G5-3 GAATCCACCATTCTTCCGAA[A/G]CTGCTTCCTACAAAACTCGA 213 132 211 S32/G3/G5-2 ATGAATTGAAGCTCTGAATA[C/T]AGAATCCACCATTCTTCCGA 214 165 211 S31/G3/G5-1 ICAATGTCTTGTTCGTTATCA[A/G]CGAAAGTTTGAATCCCCACA 215 198 211 S31/G3/G4-1 ITGTATCGGCTAGTCTGGATG[G/A]TCGCACTGGGACTCAGTGCT 216 231 211 S29/G4/G5-1 ICCTTACACTATTAACAGGCC[C/T]GTGATCTACCTGAATGCCTG 217 0 219 S29/G1/G5-1 ICCTTACACTATTAACAGGCC[C/T]GTGATCTACCTGAATGCCTG 218 33 219 S28/G2/G6-1 IGAATGTTGCTGTTATATTAC[T/C]CGTAGGTGACAAAGGGTTCA 219 66 219 S28/G3/G6-1 IGCGTAGCTTCCTCAAGACC[T/]TCCTGATGTGCGGACGCAAGC 220 99 219 S28/G3/G4-2 IAGGGGGTGGTCGCGACTGGA[T/G]CGCCCGAGCAGCGAAGC 221 132 219 S28/G3/G6-1 ITACATCTTAACAAGACC[T/]TCCTGATGTGCGGACAGCC 222 165 219 S28/G5/G6-1 ICAAGAAGCCTCTTCAGTGTC[A/C]GTCGTAGCTTCCTCAAGACC 223 198 219 S27/G3/G6-2 ICCAAGAAGATTAATGCTGG[/T]TAAAATATTGTTTCCAGTCT 224 231 219 S26/G5/G6-1 ICCAAGAAGATTAATGCTGGG[/T]TAAAATATTGTTTCCAGTCT	205	132	203	535/G4/G5-1	GGTTCGCACCATATCATGAT [C/T] GGAATGCCGCCTCAAAATGG
208 231 203 S33/G4R/G6* IGGCAGACAACAGACAGATCA [AG/CA]CATGCTTGCATTTACTCTCA 209 0 211 S33/G4r/G2* IGGCAGACAACAGACAGATCA [A/C]GCATGCTTGCATTTACTCTCA 210 33 211 S33/G5/G6-2 TCAAAGTGGTGCAATCTCGCAA [T/C]CCACTTGGGCTTGCCGTGGT 211 66 211 S33/G2R/G3-1 ACGCATGCTTGCATTTACTC [C/T]CAGTCAAACTCGCAA 212 99 211 S32/G3/G5-2 GAATCCACCATTCTTCCGAA [A/G]CTGCTTCCTACAAAACTCGA 213 132 211 S32/G3/G5-2 ATGAATTGAAGCTCTGAATA [C/T]AGAATCCACCATTCTTCCGA 214 165 211 S31/G3/G5-1 ICAATGTCTTGTTACTACA [A/G]CGAAAGTTTGAATCCCCACA 215 198 211 S31/G3/G4-1 ITGTATCGGCTAGTCTGGATG [G/A]TCGCACTGGCACTCAGTGCT 216 231 211 S29/G4/G5-1 ICCTTACACTATTAACAGGCC [C/T]GTGATCTACCTGAATGCCTG 217 0 219 S29/G1/G5-1 ICCTTACACTATTAACAGGCC [C/T]GTGATCTACCTGAATGCCTG 218 33 219 S28/G2/G6-1 IGAATGTTGCTGTATATTAC [T/C]CGTAGGTGACAAAGGGTTCA 220 99 219 S28/G3/G4-2 AGGGGGTGGTCGCGACTGGA [T/G]CGCCCGAGCAGCCCAAGC 221 132 219 S28/G3/G4-2 AGGGGGTGGTCGCGACTGGA [T/G]CGCCCGAGCAGCCGAGCAAGC 221 132 219 S28/G3/G6-1 ITACATCTTAACAAGCACTGTTTTTTTTACAACTTTTCAAACTTT 222 165 219 S28/G5/G6-1 ICAAGAAGCCTCTTCAGTGTC [A/C]GTCGTAGCTTCCTCAAGACC 223 198 219 S27/G3/G6-2 ITTCGCTTGGTAGCCGTAGCA [G/A]TATACTTTTACCGGCCCACAG 224 231 219 S26/G5/G6-1 ICCAAGAAGACTCTTCAGTGTC [A/C]GTCGTAGCTTCCTCAAGACC 224 231 219 S26/G5/G6-1 ICCAAGAAGACTTTTCAGCGGCGTAGCA [G/A]TATACTTTTACCGGCCCACAG 224 231 219 S26/G5/G6-1 ICCAAGAAGATTAATGCTTGG [/T]TAAAATATTTTTACCGGCCCACAG 224 231 219 S26/G5/G6-1 ICCAAGAAGATTAATGCTTGG [/T]TAAAATATTTTTCCAAGTCT	206	165	203	1534/G3/G5-2	GATTTTGTAGGTTGATGCAT [C/T]GTTTGATCTTTTTTTTCTCC
209 0 211 533/G4F/G2* GGCAGACAACAGACAGATCA (\(\lambda\)/C) GCATGCTTGCATTTACTCTCA 210 33 211 533/G5/G6-2 TCAAAGTGGTGCAATCGCAA (T/C) CCACTTGGGCTTGCCGTGGT 211 66 211 533/G2R/G3-1 \(\lambda\)CCATTCTTCCAATCTC (C/T) CAGTCAAACTCAGTCCCGAA 212 99 211 932/G3/G5-3 GAATCCACCATTCTTCCGAA (\(\lambda\)/G) CTGCTTCCTACAAAACTCGA 213 132 211 532/G3/G5-2 \(\lambda\)TGAATTGAAGCTCTGAATA (C/T) AGAATCCACCATTCTTCCGA 214 165 211 531/G3/G5-1 CCAATGTCTTGTTGTTATCA (\(\lambda\)/G) CGAAAGTTTGAATCCCCACA 215 198 211 S31/G3/G4-1 TGTATCGGCTAGTTGGATG (G/\(\lambda\)) TCGCACTGGCACTCAGTGCT 216 231 211 S29/G4/G5-1 TCTATTCAGCAGTCTGAGAA (GCA/CT) \(\lambda\)GGAATGCTCGGCTTCTTCAG 217 0 219 S29/G1/G5-1 CCTTACACTATTAACAGGCC (C/T) \(\lambda\)GGATCGTCGGCTTCATCAG 218 33 219 S28/G2/G6-1 GAATGTTGCTGTTATATTAC (T/C) \(\cappa\)CGTAGGTGACAAAGGGTTCA 220 99 219 S28/G3/G4-2 \(\lambda\)GGGGGGTGGTCGCGACTGGA (T/G) \(\cappa\)CGCCGAGCAAGC 221 132 219 S28/G3/G6-1 \(\tag{TACATCTTAACAAGACCATGTGT}\)CTTTTTTTTTTTTTTTAACAACTTT 222 165 219 S28/G5/G6-1 CAAGAAGCCTCTTCAGTGTC (\(\lambda\)/C (\(\lambda\)) TACATCTTAACAAGCACATG (T/C) TATATCTTTACCGGCCACAG 223 198 219 S27/G3/G6-2 \(\tag{TTCGCTTGGTAGCCGTAGCA}\)(G/\(\lambda\)) TACATCTTGAGCCGTAGCA (G/\(\lambda\)) TATACTTTTACCGGCCACAG 224 231 219 S26/G5/G6-1 CCAAGAAGACTTTAATCCTGG (/T) TAAAATATTGTTTCCAGTCT	207	198	203	S34/G2/G5-1	TGTAGGACTTGGAGAGCTTG[A/G]TAATTTACACATGCCTCTGT
210 33 211 S33/G5/G6-2 TCAAAGTGGTGCCAA [T/C]CCACTTGGGCTTGCCGTGGT 211 66 211 S33/G2R/G3-1 ACGCATGCTTGCATTTACTC[C/T]CAGTCAAACTCGCAA 212 99 211 S32/G3/G5-3 GAATCCACCATTCTTCCGAA [A/G]CTGCTTACCACAAACTCGCA 213 132 211 S32/G3/G5-2 ATGAATTGAAGCTCTGAATA [C/T]AGAATCCACCATTCTTCCGA 214 165 211 S31/G3/G5-1 CAATGTCTTGTTATCA [A/G]CGAAAGTTTGAATCCCCACA 215 198 211 S31/G3/G4-1 TGTATCGGCTAGTCTGGATG [G/A]TCGCACTGGCACTCAGTGCT 216 231 211 S29/G4/G5-1 TCTATTCAGCAGTCTGAGAA [GCA/CT]AGGATGGTCGGCTTCTTCAG 217 0 219 S29/G1/G5-1 CCTTACACTATTAACAGGCC [C/T]GTGATCTACCTGAATGCCTG 218 33 219 S28/G2/G6-1 GAATGTTGCTGTTATATTAC [T/C]CGTAGGTGACAAAGGGTTCA 220 99 219 S28/G3/G4-2 AGGGGGTGGTCGCGACTGGA [T/G]CGCCCGAGCAGCCGAGCAAGC 221 132 219 S28/G3/G4-2 TACATCTTAACAAGCACTGTGTTTTTAACCTTTTTTTCAAACTTT 222 165 219 S28/G5/G6-1 CCAAGAAGCCTCTCAGTGTC [A/C]GTCGTAGCTTCCTCAAGACC 223 198 219 S27/G3/G6-2 TCCGCTTGGTAGCCGTAGCA [G/A]TATACTTTTACCAGGCCACAG 224 231 219 S26/G5/G6-1 CCAAGAAGCATTAATCCTGG [/T]TAAAATATTGTTTCCAGTCT	208	231	203	S33/G4R/G6*	GGCAGACAACAGACAGATCA AG/CA CATGCTTGCATTTACTCTCA
211 66 211 S33/G2R/G3-1 ACGCATGCTTGCATTTACTC[C/T]CAGTCAAACTCAGTCCCGAA 212 99 211 S32/G3/G5-3 GAATCCACCATTCTTCCGAA[A/G]CTGCTTCCTACAAAACTCGA 213 132 211 S32/G3/G5-2 ATGAATTGAAGCTCTGAATA[C/T]AGAATCCACCATTCTTCCGA 214 165 211 S31/G3/G5-1 CAATGTCTTGTTCGTTATCA[A/G]CGAAAGTTTGAATCCCCACA 215 198 211 S31/G3/G4-1 ITGTATCGGCTAGTTCTGGATG[G/A]TCGCACTGGCACTCAGTGCT 216 231 211 S29/G4/G5-1 ICCTTACACTATTAACAGGCC[C/T]AGGATGGTCGGCTTCTTCAG 217 0 219 S29/G1/G5-1 ICCTTACACTATTAACAGGCC[C/T]GTGATCTACCTGAATGCCTG 218 33 219 S28/G2/G6-1 IGAATGTTGCTGTTATATTAC[T/C]CGTAGGTGACAAAGGGTTCA 219 66 219 S28/G3/G6-1 IGTCGTAGCTTCCTCAAGACC[T/]TCCTGATGTGCGGACGCAAGC 220 99 219 S28/G3/G4-2 IACGGGGTGGTCGCGACTGGA[T/G]CGCCCGAGCAGCCAAGC 221 132 219 S28/G3/G6-1 ITACATCTTAACAAGCACATG[TG/TTT]TAACCTTTTATTCAAACTTT 222 165 219 S28/G5/G6-1 ICAAGAAGCCTCTTCAGTGTC[A/C]GTCGTAGCTTCCTCAAGACC 223 198 219 S27/G3/G6-2 ITTCGCTTGGTAGCCGTAGCA[G/A]TATACTTTTACCGGCCACAG 224 231 219 S26/G5/G6-1 ICCAAGAAGATTAATGCTGG[/T]TAAAAATATTTTTTCCAGTCTCT	209	0	211	S33/G4r/G2*	GGCAGACAACAGACAGATCA (A/C) GCATGCTTGCATTTACTCTCA
212 99 211 932/G3/G5-3 GAATCCACCATTCTTCCGAA[A/G]CTGCTTCCTACAAAACTCGA 213 132 211 532/G3/G5-2 ATGAATTGAAGCTCTGAATA[C/T]AGAATCCACCATTCTTCCGA 214 165 211 531/G3/G5-1 CAATGTCTTGTTCGTTATCA[A/G]CGAAAGTTTGAATCCCCACA 215 198 211 531/G3/G4-1 TGTATCGGCTAGTCTGGATG[G/A]TCGCACTGGCACTCAGTGCT 216 231 211 529/G4/G5-1 TCTATTCAGCAGTCTGAGAA[GCA/CT]AGGATGGTCGGCTTCTTCAG 217 0 219 529/G1/G5-1 CCTTACACTATTAACAGGCC[C/T]GTGATCTACCTGAATGCCTG 218 33 219 528/G2/G6-1 GAATGTTGCTGTTATATTAC[T/C]CGTAGGTGACAAAGGGTTCA 220 99 219 528/G3/G4-2 AGGGGGTGGTCGCGACTGGA[T/G]CGCCCGAGCAGCGCAAGC 221 132 219 528/G3/G4-2 AGGGGGTGGTCGCGACTGGA[T/G]CGCCCGAGCAGCCGAACCC 221 132 219 528/G3/G6-1 CAAGAAGCCTCTTCAGTGTC[A/C]GTCGTAGCTTCCTCAAGACC 223 198 219 527/G3/G6-2 TTCGCTTGGTAGCCGTAGCA[G/A]TATACTTTTACCGGCCACAG 224 231 219 526/G5/G6-1 CCAAGAAGATTAATGCTGG[/T]TAAAATATTGTTTCCAGTCT	210	33	211	S33/G5/G6-2	TCAAAGTGGTGCAATCGCAA (T/C) CCACTTGGGCTTGCCGTGGT
132 211 532/G3/G5-2 ATGAATTGAAGCTCTGAATA[C/T]AGAATCCACCATTCTCCGA 214	211	66	211	S33/G2R/G3-1	ACGCATGCTTGCATTTACTC [C/T] CAGTCAAACTCAGTCCCGAA
214 165 211 S31/G3/G5-1 CAATGTCTTGTTATCA (A/G) CGAAAGTTTGAATCCCCACA 215 198 211 S31/G3/G4-1 TGTATCGGCTAGTCTGGATG (G/A) TCGCACTGGCACTCAGTGCT 216 231 211 S29/G4/G5-1 TCTATTCAGCAGTCTGAGAA (GCA/CT) AGGATGGTCGGCTTCTTCAG 217 0 219 S29/G1/G5-1 CCTTACACTATTAACAGGCC (C/T) GTGATCTACCTGAATGCCTG 218 33 219 S28/G2/G6-1 GGAATGTTGCTGTTATATTAC (T/C) CGTAGGTGACAAAGGGTTCA 219 66 219 S28/G3/G6-1 GTCGTAGCTTCCTCAAGACC (T/) TCCTGATGTGCGGACGCTAA 220 99 219 S28/G3/G4-2 AGGGGGTGGTCGCGACTGGA (T/G) CGCCCGAGCAGCGAGCAAGC 221 132 219 S28/G3/G3-1 TACATCTTAACAAGCACATG (TG/TTT) TAACCTTTTATTCAAACTTT 222 165 219 S28/G5/G6-1 ICAAGAAGCCTCTTCAGTGTC (A/C) GTCGTAGCTTCCTCAAGACC 231 198 219 S27/G3/G6-2 TTCGCTTGGTAGCCGTAGCA (G/A) TATACTTTTACCGGCCACAG 224 231 219 S26/G5/G6-1 CCAAGAAAGATTAATGCTGG (/T) TAAAATATTGTTTCCAGTCT	212	99	211	932/G3/G5-3	GAATCCACCATTCTTCCGAA (A/G) CTGCTTCCTACAAAACTCGA
215 198 211 S31/G3/G4-1 ITGTATCGGCTAGTCTGGATG[G/A] TCGCACTGGCACTCAGTGCT 216 231 211 S29/G4/G5-1 ITCTATTCAGCAGTCTGAGAA [GCA/CT] AGGATGGTCGGCTTCTTCAG 217 0 219 S29/G1/G5-1 ICCTTACACTATTAACAGGCC [C/T]GTGATCTACCTGAATGCCTG 218 33 219 S28/G2/G6-1 IGAATGTTGCTGTTATATTAC [T/C]CGTAGGTGACAAAGGGTTCA 219 66 219 S28/G3/G6-1 IGTCGTAGCTTCCTCAAGACC [T/]TCCTGATGTGCGACCAGCGAGCAAGC 220 99 219 S28/G3/G4-2 AGGGGGTGGTCGCGACTGGA [T/G]CGCCCGAGCAGCGAGCAAGC 221 132 219 S28/G1/G3-1 ITACATCTTAACAAGCACATG [TG/TTT]TAACCTTTTATTCAAACTTT 222 165 219 S28/G5/G6-1 ICAAGAAGCCTCTTCAGTGC [A/C]GTCGTAGCTTCCTCAAGACC 223 198 219 S27/G3/G6-2 ITCGCTTGGTAGCCGTAGCA [G/A]TATACTTTTACCGGCCACAG 224 231 219 S26/G5/G6-1 ICCAAGAAAGATTAATGCTGG [/T]TAAAATATTGTTTCCAGTCT	213	132	211	532/G3/G5-2	ATGAATTGAAGCTCTGAATA [C/T] AGAATCCACCATTCTTCCGA
216 231 211 S29/G4/G5-1	214	165	211	IS31/G3/G5-1	
217 0 219 S29/G1/G5-1 ICCTTACACTATTAACAGGCC [C/T]GTGATCTACCTGAATGCCTG 218 33 219 S28/G2/G6-1 IGAATGTTGCTGTTATATTAC [T/C]CGTAGGTGACAAAGGGTTCA 219 66 219 S28/G3/G6-1 IGTCGTAGCTTCCTCAAGACC [T/]TCCTGATGTGCGGACGCGAGCCAAGC 220 99 219 IS28/G3/G4-2 IAGGGGGTGGTCGCGACTGGA[T/G]CGCCGAGCAGCCAAGC 221 132 219 S28/G1/G3-1 ITACATCTTAACAAGCACATG[TG/TTT]TAACCTTTTATTCAAACTTT 222 165 219 S28/G5/G6-1 ICAAGAAGCCTCTTCAGTGTC[A/C]GTCGTAGCTTCCTCAAGACC 223 198 219 S27/G3/G6-2 ITTCGCTTGGTAGCCGTAGCA[G/A]TATACTTTTACCGGCCACAG 224 231 219 S26/G5/G6-1 ICCAAGAAAGATTAATGCTGG [/T]TAAAAATATTGTTTCCAGTCT	215	198	211	S31/G3/G4-1	
218 33 219 S28/G2/G6-1 GAATGTTGCTGTTATATTAC [T/C]CGTAGGTGACAAAGGGTTCA 219 66 219 S28/G3/G6-1 GTCGTAGCTTCCTCAAGACC [T/]TCCTGATGTGCGGACGCTAA 220 99 219 S28/G3/G4-2 AGGGGGTGGTCGCGACTGGA [T/G]CGCCCGAGCAGCGAGCAAGC 221 132 219 S28/G1/G3-1 ITACATCTTAACAAGCACATG [TG/TTT]TAACCTTTTATCAAACTTT 222 165 219 S28/G5/G6-1 ICAAGAAGCCTCTTCAGTGTC [A/C]GTCGTAGCTTCCTCAAGACC 223 198 219 S27/G3/G6-2 ITTCGCTTGGTAGCCGTAGCA [G/A]TATACTTTTACCGGCCACAG 224 231 219 S26/G5/G6-1 ICCAAGAAAGATTAATGCTGG [/T]TAAAAATATTGTTTCCAGTCT	216	231	211	IS29/G4/G5-1	
219 66 219 IS28/G3/G6-1 GTCGTAGCTTCCTCAAGACC[T/]TCCTGATGTGCGGACGCTAA 220 99 219 IS28/G3/G4-2 AGGGGGTGGTCGCGACTGGA[T/G]CGCCCGAGCAGCGAGCAAGC 221 132 219 S28/G1/G3-1 ITACATCTTAACAAGCACATG[TG/TTT]TAACCTTTTATTCAAACTTT 222 165 219 S28/G5/G6-1 ICAAGAAGCCTCTTCAGTGTC[A/C]GTCGTAGCTTCCTCAAGACC 223 198 219 S27/G3/G6-2 ITTCGCTTGGTAGCCGTAGCA[G/A]TATACTTTTACCGGCCACAG 224 231 219 S26/G5/G6-1 ICCAAGAAAGATTAATGCTGG[/T]TAAAAATATTGTTTCCAGTCT	217	0	219	IS29/G1/G5-1	CCTTACACTATTAACAGGCC C/T GTGATCTACCTGAATGCCTG
220 99 219 IS28/G3/G4-2 AGGGGGTGGTCGCGACTGGA(T/G)CGCCCGAGCAGCGAGCAAGC 221 132 219 S28/G1/G3-1 ITACATCTTAACAAGCACATG[TG/TTT]TAACCTTTTATTCAAACTTT 222 165 219 S28/G5/G6-1 ICAAGAAGCCTCTTCAGTGTC[A/C]GTCGTAGCTTCCTCAAGACC 223 198 219 S27/G3/G6-2 ITTCGCTTGGTAGCCGTAGCA [G/A]TATACTTTTACCGGCCACAG 224 231 219 S26/G5/G6-1 ICCAAGAAAGATTAATGCTGG [/T]TAAAAATATTGTTTCCAGTCT	218	33	219	S28/G2/G6-1	GAATGTTGCTGTTATATTAC(T/C)CGTAGGTGACAAAGGGTTCA
221 132 219 \$28/\$G\$1/\$G\$3-1 TACATCTTAACAAGCACATG[TG/TTT]TAACCTTTTATTCAAACTTT 222 165 219 \$28/\$G\$5/\$G\$6-1 CAAGAAGCCTCTTCAGTGTC[A/C]GTCGTAGCTTCCTCAAGACC 223 198 219 \$27/\$G\$3/\$G\$6-2 TTCGCTTGGTAGCCGTAGCA[G/A]TATACTTTTTACCGGCCACAG 224 231 219 \$26/\$G\$5/\$G\$6-1 CCAAGAAAGATTAATGCTGG[/T]TAAAATATTGTTTCCAGTCT	219	66	219	1528/G3/G6-1	
222 165 219 \$28/\$\text{G5}/\$\text{G6-1}\$ CAAGAAGCCTCTTCAGTGTC [A/C]GTCGTAGCTTCCTCAAGACC 223 198 219 \$27/\$\text{G3}/\$\text{G6-2}\$ TTCGCTTGGTAGCCGTAGCA [G/A]TATACTTTTACCGGCCACAG 224 231 219 \$26/\$\text{G5}/\$\text{G6-1}\$ CCAAGAAAGATTAATGCTGG [/T]TAAAATATTGTTTCCAGTCT	220	99	219	1528/G3/G4-2	
223 198 219 S27/G3/G6-2 ITTCCCTTGGTAGCCGTAGCA(G/A)TATACTTTTACCGGCCACAG 224 231 219 S26/G5/G6-1 CCAAGAAAGATTAATGCTGG[/T]TAAAATATTGTTTCCAGTCT	221	132	219	S28/G1/G3-1	TACATCTTAACAAGCACATG TG/TTT TAACCTTTTATTCAAACTTT
224 231 219 S26/G5/G6-1 CCAAGAAAGATTAATGCTGG[/T]TAAAATATTGTTTCCAGTCT	222	165	219	S28/G5/G6-1	CAAGAAGCCTCTTCAGTGTC A/C GTCGTAGCTTCCTCAAGACC
	223	198	219	S27/G3/G6-2	ITTCCCTTGGTAGCCGTAGCA (G/A) TATACTTTTACCGGCCACAG
	224	231	219	S26/G5/G6-1	
225 0 227 S26/G4/G5-2 SATGCTGGTAAATATTGTTT (C) CAGTCTTTCACAAAGTGTGT	225	0	227	S26/G4/G5-2	ATGCTGGTAAAATATTGTTT /C CAGTCTTTCACAAAGTGTGT

226	33	227	S23/G5/G3-1	TTAAGTGAAGATGCCCAAAC [C/G]GTTAAACTTTCCATGGAACT
227	66	227	S23/G1/G5-1	CGGGTCTGTATGCGAGTGTT (G/A)TGGTGGTGAACTGGTGAATT
228	9 9		S23/G4/G5-2	TTCCAGTCGGATGAACTGGA [T/G]GTTCGTCATCCACTCGTCAC
229	132		S22/G5/G6-1	CAGATTGGTGTCGTTTACTA (A/G) AATTCAGTTCTGTCCATTTG
230	165	227	S21/G2/G5-1	TGGCTTGGTTGACCTTGAGG[C/G]CCACACACTATCTAGTACGT
231	198	227	S20/G5/G6-1	GCCGCCGAGAGCGAGGCATA (G/T) GCGCATGTGCATGTGCGTGC
232	231	227	S19/G5/G6-2	AGTACATGGCGAGCGTTGTA [G/C] CAGCTGCTTAGGTGATGTGG
233	0	235	S17/G3/G6-1	GGTTCTAAACATAGCTCGTC [C/A]ATTCATGATTCATCTCGAGC
234	33	235	S17/G4/G6-1	CTATTTCC&AGCTAACAACC [C/G] CTCTTGGTCCCAACATCCTG
235	66	235	517/G3/G7-3	AAGCCTCATGTGCAGATTCA (TC/AG) GAACACACAACGTCAGCCAT
236	99	235	S16/G1/G2-1	GTGTGTAGCTTCATTCGCAA [A/T]GTTTGAACAGCCTCTGCAAG
237	132	235	S16/G4/G6-1	TCAGCAAGCCTCCAAGGCTC [C/A]AATGGTCCAGTTACTTGGTT
238	165	235	S16/G2/G7-2	GTTGACCAGCTGTGATTTCG[G/A]TGTATTCCACGACCACGAGT
239	198	235	516/G1/G7-1b	CTTAATTGTACACAGTGCTT [C/T] CGTAAACCTAGAGCLGACCA
240	231	235	S16/G1/G7-1c	CTTAATTGTACACAGTGCTT [C/T] CGTAAACCTAGAGLGGACCA
241	0	243	S16/G6/G7-1a+	GTGCTECCGTAAACCTAGAG (TT/CT) GACCAGCTGTGATTTCGATG
242	33	243	S16/G6/G7-1b*	GTGCTTTCGTAAACCTAGAG (CT/TG)GACCAGCTGTGATTTCGATG
243	66	243	S14/G5/G7-2	CGAAAGAGCGAGATATATCG [A/G]TCGATCGATGAGCAAGTATA
244	99	243	S14/G5/G6-1	GCTCAGCTGCCGGAGTACGT[A/T]GGCTTGCTCTCCGGCCGGCC
245	132	243	S13/G2/G6-1	TTGGTAATTTCAGAGCTAGA [C / G] AACTTACTGTGGTACACGCC
246	165	243	S13/G5/G6-1	TTTCACAACTCAACTGATTG[A/T]CTTGCTTTGATGTGGATTCT
247	198	243	S12/G2/G5-3a	CGTAATTACTTTGTTACESC [TA/C]AGTAATTTTATATATATCCT
248	231	243	S12/G2/G5-3b	CGTAATTACTTTGTTACacc[TA/C]AGTAATTTTATATATATCCT
249	0	251	S12/G1/G5-3	CTTTACTGATTGGGTTACAA [A/G]AGGTTATTTCTTATTCAGGC
250	33	251	S12/G2/G4-3a	ATCCGTARTTACTTTGTTAC TA/AC CLAAGTARTTTTATATATAT
251	66	251	S10/G4/G6-2	AATTTGGGAAAATCAATGCA (GAA/CAC) ATCAGTGATTAATCCACATA
252	99	251	S10/G4/G6-1	AGCGACAGGGATGTCGAGCA (G/T)CTACGGAAGGCAATAATGAG
253	132	251	S08/G3/G6-1*	GCACGTCGTTGGTGAAGAAG (AC/CA)GCGGTACGGGTGCTTGTCGA
254	165	251	S08/G1/G4-1*	GCACGTCGTTGGTGAAGAAG (A/C)AGCGGTACGGGTGCTTGTCG
255	198	251	S08/G3/G5-1	AGGTACACGGGGAAGTCGGA (G/T)TGGTTCTTCACCACCACCGC
256	231	251	S08/G5/G6-1	GTCCCAGATCAGGTCCACGT (T/C) CGAGCTCGCTGTTCCCGCTT
257	0	259	S06/G2/G3-2	ACGGTGAGGAGTGGCACATG (A/C)GATGGAAAGTTCCTGTAGAC
258	33	259	S06/G2/G3-1	INAACCAAACCCTGACTATTA (T/C) AGGTAGATTAGACTAGACAC
259	66		S03/G2/G6-1	ATATCCATGTTGTCGCCTGC[/TG]TGTGCGCTTGCTTGCCGCTA
260	99	259	S02/G2/G5-2	GTGTGGAATGACCATCTCGT [G/C]GTGATGCCAGCATGCTGTTA
261	132	259	502/G3/G7-1b	CAACGTGCAATAATAGAACA [T/G]GTGGTGTTTGTTAAGAAAGA
262	165	259	S02/G6/G7-1	CACGGCAGTTGGCAGTGTGG (A/) AAGGACTATCTCGTGGTGAT
263	198	259	S01/G4/G7-3	GCTAGAGCAAGAGTCAACAC (G/A) CGCGCGCATCACGCATGCCA
264	231	259	S01/G3/G7-2	AGCCAGGTTCTAACAGCTAG [C/A]GCAAGAGTCAACACGCGCGC

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sequence
CTTACATGCGTGGAAAGAGA (G/)TCCTCGAGATCAACCCACGA
GATGCTAAAAAGCAAGCTTA [C/T] ATGCGTGGAGTGAGAGTCCT
AGCAAGCTTACATGCGTGGA (GT/AA) GAGAGTCCTCGAGATCAACC
CAAACTCTCTCAGATGCTAA (A/G) AAGCAAGCTTACATGCGTGG
CARACTETETETETETETETETETETETETETETETETETETE
CCTTGATCTCTCAAGTAATC (A/G)TCTCACCGGAAGATCCCTGA
TCTCGATCTGACATCTCTCA (A/T) CGTCCTTGATCTCTCAAGTA
TCGTCGCAATCTAACCAAAA (C/G)TTAAAAACGCTAACGGTGTG
TCGTCGCAATCTAACCAAAA(C/G)TTAAAAACGCTAACAGTGTG
TCATCGCAATCTAACCAAAA (C/G)TTAAAAACGCTAACGGTGTG
TCATCGCAATCTAACCAAAA [C/G]TTAAAAACGCTAACAGTGTG
ACCATCCATTAAACTGTATC (A/G)TCGCAATCTAACCAAAAGTT
TARAGCAAAGAGAGTCTTAC(C/A)GTCTGCTGCATGATGTACCC
TAAAGCAAAGAGTCTTAC(C/A)GTCTGCTGCATGATATACCC
CTACTGATAGTGAACCACCC(A/C)ATCCCCAAATTTAAAGCAAA
AACTTTTCGATTTCAATATA(G/T)CAACTACTGATAGTGAACCA
TAACTCTACAACATTCACAA(C/G)CTGGGCCAACATATTTAAAC
GGGATATAGCAAANTTGAAC (A/C) ATATCAATTGTTGACGGAAC
· ·
CCAAATAGGTTCTGAGCCTT(C/T)ACTTTCTCCACCATCTCATT
CTCCACCATCTCATTGCTCA(T/A)TCCAGCAAAAAACATCTCTG
CTCCACCATCTCATTGCTCA [G/A]TCCAGCAAAAAACATTTCGG
CTCCACCATCTCATTGCTCA [G/A] TCCAGCAAAAAACATCTCGG
CTCCACCATCTCATTGCTCA [G/A] TCCAGCAAAAACATTTCTG
CTCCACCATCTCATTGCTCA (G/A) TCCAGCAAAAAACATCTCTG
CCARATAGGTTCTGAGCCTT[T/C]ACTTTCTGCAGCATCTCATT
CCAMINOSTICIONOCCITTICINA
ATCCTATTGGTAGTAACACA (G/A) ATTGAGTTAATGTTGCAGGG
AGGCAAAGCGGTAGTTGCAA[G/A]ACTGCTTCTCACGAGGTAAT
TCCTCAGAGGCAAAGCGGTA(T/G)TTGCAAAACTGCTTCTCACG
TCCTCAGAGGCAAAGCGGTA(1/0)110c1344Fc13c111c1ag
CTTCCCCCCCTCAATACCTC (T/G)TTCAAAAGTGAAAAGTGCAG
TOTANGATCCTCTCTCTC(CT/TA)TGTCAACTCTCCCCCCCCC
TGTAAATCCTCTCTCTCTC(CT/TA)TGTCAACTCTCCCCCCCCCCCCCCCCCCCCCCCCCCC
TGTAAATCCTCTCTCTCTCTCTCTATTGTCAACTCTCCCCCCCC
TTTCAAAGTGAAAAGTGCA [G/A]AAACTCATTTATTTATGTTT
TTACAAAAGTGAAAAGTGCA(G/A)AAACTCATTTATTTATGTTT
GTTCARAAGTGAAAAGTGCA (G/A) AAACTCATTTATTTATGTTT
GTACAAAAGTGAAAAGTGCA[G/A]AAACTCATTTATTTATGTTT
AAAGTTCATTACGATGATCT (A/G) ACCCTGCAGTCATCCATGGA
AAAGTTCATTACGATGATCT (A/G)ACCCTGCAGTCATCCATAGA CCAGCTTCAATGTCTGCATG(C/A)TTGTGTCGATGCCAAAGTTC

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N1/6D1/N2-1A	ATTTTGTTTTGTTCTGTC(G/C)GGTCAGGTCAGAACAAAGTT
N1/6D1/N2-1B	ATTTTGTTTTGTTTGTC [G/C] GGTCAGGTCAGAACAAAGGT
	Ar- •
N1-6F11-N2	TAGGACAGGCAAACAATCTA (C/A) GCGGTCAAAATCCGATTTCG
86-6F11-N2-1	GAGGATGAGTATATCGCTTT (A/G)TGTCTCATGCTTCTTGCTCG
86-6F11-N2-2	GATTTCGACCGCAGTCTCAC [G/T] GAGGATGAGTATATCGCTTT
N1-6H5-N2	AAACCAGAGCCACCTCCTTA(C/)CCACCTCATCGTTTCCTTTC
L4-8A2-L6-1	ACCGAAACCAATCTCCCAAG [T/]AAAGCTTATTCAGGAGCTTC
L4-8A2-L6-2	AGCAGTGCAACAGCTCCTGC (A/T) AAGTCCCTGACGTACGAGGA
L4-8A2-L6-3	AGCTTCAATGGCGGATGGGC [T/C] CTTCTAACCTCTGTTCTAAG
N1/8B5/N2	ACTCAAAAAACGATACCTC[G/C]GCCGTCTCTCGCCGTCTCGC
85-8B5-86-1	AAACACTAAGTGTCNCTCTC (T/C) AAACTAGTGTGCAAAGCTCA
N1/8D4/N2-1A	CAGGAGACAGTTACAGTCCC(/A)CAGGACTCGCAAGGATCTCGAA
N1/8D4/N2-1B	CAGGAGACAGTTACAGTCCC[/A]CAGAGTCGCAAGGACTCGAAC
86/8D4/N1-1	ATCTCGAACTTCACATCTGC [G/A] TTGAGTTCTGCTGAGAGGCT
85-8D4-86-1A	AGTTACAGTCCCACAGAGTC(T/G)CAAGGATCTCGAACTTCACT
85-8D4-86-1B	ACTTACAGTCCCACAGAGTC[T/G]CAAGGATCTCGAACTTCACA
85-8D4-86-2	CTGATCTTGAAGGAGAGACC [A/G] CCACAAGGTTCCATCCTATG
85/8H11/86	AGTGCNAGGCTCAGTTGGAT [G/T]ATTAGGGTGTCAGTAAATCA
N2/10B8/N3-1A	AAGGTATTCCATTGGTATAC [\(\lambda / C \) TCCAACTTTTCTCCAGTTCT
N2/10B8/N3-1B	AAGGTATTCCATTGGTATAC (A/C)TCCAACTTTTCTCCAGTCCT
N2/10B8/N3-2A	ACATCCAACTTTTCTCCAGT[T/C]CTTTATTCTATCCTGATTTG
N2/10B8/N3-2B	ACCTCCAACTTTTCTCCAGT[T/C]CTTTATTCTATCCTGATTTG
85-1088-86	NAOGTCCATGATGACAA [T/A] AAAGGTATTCCACATGTCAA
	•
85-10B9-86	GACCTTCTTGGGAAAGAAAG [T/C]TGTAACCGCGTCGAGATTCG
N2/10B9/N3A	CAGATCGGTACTTTCAACCA(G/T)TCTCCTTCTCTGTCTCCACT
N2/10B9/N3B	AGATCCGGTACTTTCAACCA[G/T]TCTCCTTCTCTGTCTCCACT
N2/10B9/N3C	AGAGTCGGTACTTTCAACCA(G/T)TCTCCTTCTCTGTCTCCACT
N2/10B9/N3D	CAGACCGGTACTTTCAACCA [G/T]TCTCCTTCTCTGTCTCCACT
86/10B9/N3	TATTCAACCAGTCTCCTTCT [T/C]TGTCTCCACTATGTCGTTAG
85/10B9/N3	ACTGAGTTTGGTCTGTCTCT [A/G]TGTGTGTGTGTTGTAGTA
L6-10C8-N2	CACTTTCTTCGTGGCTAAAT [G/T] CTTCGGCCGAGCCGGTCTCA
L5-10C8-N2	ATAGAAACCGCCGATGCTCA [G/A] GGACACGCCACCGTCTTCGT
L6-10D2-N1	GTTATCATCAGTACCGGTAT (C/T)AACCCCAAGGCTAATTCTTA
L5-10D2-N2	GTGTTGGGTATCTACGGACT[C/G]ATCATCGCTGTTATCATCAG
85-10D2-L6	TTGGGTATCTACGGACTGAT (C/T) ATCGCTGTTATCATCAGTAC
85-10D2-L6	TAGGGTATCTACGGACTGAT (C/T) ATCGCTGTTATCATCAGTAC
N1/10E12/N2-1	GGAATTCAATACTCGCCAAC [G/T]TCTTCATTGCTGTCGTCGGC

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N1/10E12/N2-2	TCCTTACGCCTTCAAGCGCA[C/G]CGGCTGGCTCATGGGTGTCC
N1/10F4/N2	TGTATCTATGCGGTGGCTGC [G/C] GTCTCCGTTCGCGCCAGTAC
L3-10F4-L6	CCGTCTCCGTTCGCGCCACT (G/A) CCGCCGGTTACCATCTCACT
L2-10F4-L6A	GTTACAATCTCACTGCCTCC [A/T] CCTCCGCTCCTATCGAGGAA
L2-10F4-L6B	GTTACCATCTCACTGCCTCC [A/T] CCTCCGCTCCTATCGAGGAA
L1/10F4/N2	GCGCCAGTACCGCCGGTTAC [C/A] ATCTTACTGCCTTCACGTCC
L1/10F4/N1-1	CTGACTCGTGCGGTGGCTGC [C/G]GTCTCCGTTCGCGCCAGTAC
L1/10F4/N1-2	CCTCCACGTCCGCTCCTATC [G/A] AGGAATCGATCGTGTCTCAC
85/10F4/N2	GCGCCAGTACCGCCGGTTAC [G/A] ATCTTAATGCCTTCACGTTC
N1-10F4-L4	GATATGATCACTTCCGCTGA[A/G]ACAGATGTTGTCGTCGTTGG
NI-IUF4-L4	CALL COLLEGE C
85/10F9/N2-1	CGGTACTGCGAAAGCTGGAG[C/G]ATCAACTTGGAATTCCACAA
85/10F9/L3-1	CGGTACTGCGAAAGCTGGAG[C/G]ATCAACTTGGAATTCCACAT
85-10F9-N1-1	ACGGTATTGTTTCTTCAGGG [T/G]TCTCGACAAACCTGAAACGG
L5-10F9-N2	GATCAACTTGGAATTCCACA(T/A)CTTCAGAAACTTCGATGTGG
L1-10F9-L4A	CAAACCTGAAACGGTACTGC [G/A] AAAGCTGGAGGATCAACTTG
L1-10F9-L4B	CAAACCTGAAACGGTACTGC [G/A] AAAGCTGGAGGATCAAGTTG
85/10P9/N1-2	AACTTGGAATTCCACAACTT [G/C] AGAAACTTCGATGTGGTGCC
L2-10F9-L4	TCAGGGGTCTCGACAAACCT (A/G) AAACGGTACTGCAAAAGCTG
L3-10F9-L4-1A	TGCGAAAGCTGGAGGATCAA [C/G]TTGGAATTCCACATCGTCAG
L3-10F9-L4-1B	TGCAAAAGCTGGAGGATCAA [C/G]TTGGAATTCCACATCGTCAG
L5-10F9-L6	TCAACTTGGAATTCCACATC[T/G]TCAGAAACTTCGATGTGGTG
85/10F9/L1-1	CAATTGCGGATGTCTTCATC [G/A] TCAACCCAAAACGGTATTGT
86-10F12-L3	AAAAGTGCTATTGTTCAGGT [G/C] GATGCTGCTCCGTTCAAGCA
85-10F12-86	AAACCAGGGTCCTTGATGTG[T/]GTCTACAACGCTTCCAACAA
L1-10H6-L6	GCGGTGGTAACAGGCGCTCA (G/T) TTTCTCGTCAAAAGCCACGG
85-10H6-L6	TTTCTCGTCAAAAGCCACGG (A/G) TTCAAGAACGTGCTCTTCTT
85-10H6-86	GTCAAAAGCCACGGATTCAA (G/A) AACGTGCTCTTCTTGCGCCT
	·
L4-11A3-L1	TTGAAGGAGGTTGGTACACA (C/G) TTCTTCGAGCTACCGGAGAC
85-11B7-86	AANACCCTGAGCTCATGCCT(C/T)TGACCCATGTTCTTGCCACC
85-11C4-86	TTTGGGACCGTTGGAGTTGC [A/G]TCTGCGGCTATGACGGTGGA
85-11D4-86-1	GTTGTGGTAGCGACTGCGGG (G/A) CAGAGCCGGTGCGAGCCTGG
85-11D4-86-2	AATCTTTGCCATTGCTGTCA(A/G)TATCTTCGTCAGCTTCAGCT
85-11D11-86	ATGCTCAGCACCATCGCCAC [T/C] GGTTTGGCGATTGATGCTTA
N2-11D11-N3	GACAACGCTGGTGGTATTGC [C/T] GAAATGGCTGGAATGAGCCA
86-11D11-N3	GCTGCTCTAGGGATGCTCAG[C/T]ACCATCGCCACCGGTTTGGC
N2-11E3-N8	GAGAAAGTGCTTGTGGAGAT[C/T]TACAAGTCCATACTGATGGC
86-11E3-N2A	AATGCTTGTGGAGATTTACA (G/A) GTCCATACTGATGGCGCAGG
86-11E3-N2B	AGTGCTTGTGGAGATTTACA (G/A) GTCCATACTGATGGCGCAGG

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N1/11F12/N3	TGGTTCTTTGGTGGGTTGCA [A/G]TTTCAGCTAGAGCATCATCT
85-11F12-B6	AATGATTGGTTTGAGAAGCA (T/A) ACAGCTGGTACGCTTGATAT
85-11F7-86	GATAGGGCGAAGAGAGGGAA (G/A) AGTCCTGAGAGGAAAGAGAT
85-11H2-86-1	GGAGAAACCCTCTCCGACTT (C/T) CTCTCTCTCCACAAAGACAC
85-11H2-86-2	CTCTCTCCACAAGACAC [A/C] GCTTTCTCCATGACCTTCGG
85-11H5-86-1	TCTCTGACGTCATGAAAGCT (C/A)ATGGCAAAATTGCTGATGGA
85-11H6-86-1	GTTATCGATCGCGTGGTCCG(T/C)GAAACCCAAAATNCACCTTT
85/12B6/N3	CGTCAGCCTTCTTCCGCCGC (A/C)GTCGTCCTCCGCAACCGTGC
L3/12B11/N2-1	GCGGATCCTATATTGGGTCT(C/T)GATGGATTGTTTCTATCCCG
L3/12B11/N2-1B	GCGGATCCTATATTGGGTCT (C/T) GATGGATTGTTTCTATTCCG
L3/12B11/N2-2A	TCTCGATGGATTGTTTCTAT (C/T) CCGCAAATTGACATACTCCT
L3/12B11/N2-2B	TCTTGATGGATTGTTTCTAT[C/T]CCGCAAATTGACATACTCCT
L3/12B11/N2-2C	TCTCGATGGATTGTTTCTAT [C/T] CCGCAAATTGACATTTCCCT
L3/12B11/N2-2D	TCTTGATGGATTGTTTCTAT [C/T] CCGCAAATTGACATTTCCCT
L3/12B11/N2-3	TATTCATGCCAACAATCTGT[C/G]GCGGATCCTATATTGGGTCT
	TATTCATGGCAAGAATCTGT [G/T] GCCGATCCTATATTGGGTCT
L2-12P11-N2-1	TATCCTGCAAATTGACATTT[T/C]CCTTCAGGTTCTAGAAGCTG
L2-12B11-N2-2	TCAGGTTTACCTCTATATAT [T/] ATATTTCATGGTATGAAGGT
86-12B11-85	TEAGOTTIACCTC INTERIOR (17) FRIENT CANADATTICAL
06/1206/32	TGTCTCTTCCGTCAGCCTTC[C/T]TCCGCCGCAGTCGTCCTCCG
86/1286/N3	CGTCAGCCTTCTTCCGCCGC [A/C] GTCGTCCTCCGCAACCGTGC
85-1286-N3	COLCADO I I COLCADO COMO COLCADO COMO COLCADO
06 1202 06	CGAGAACAGAAGAGAGAGA(C/)TGGAACACGTCGGACAGTAC
85-12C2-86	CONORMONOMONOMONOCO
	TGNCACAACGAAGGTTTTGG [C/T] GGAGGTAAATGCCGTGGGTT
L3-12C7-L5	TGNCACAACGAAGGTTTTGG(C/T)GGAGGTAATTGCCG33GGT.
	ACGGGTCCTAGCGCCATGGC[C/T]ATTTTCCTCACCGTTTCTGG
L6-12C11-N2	ACCOSTCCTAGEGECATGGE (C/T)ATTTTCCTCACCGTTCTGG
	TOTAL CONTROL OF THE
N1-12D10-L6	TTGGGCTTTCGGTGGTATGA (T/)CTTCGTCCTCGTCTATTGCA
N1,2,3/12E10/L1	TACCACGGTCGTACTGGTCG (A/)TGTCTGGAACGTCACCAAGC
L2-12E10-L3	CACAAAGGTCTGCCTCACAA (G/T) TTCTACCACGGTCGTACTGG
85-12F4-86-1	TCCTTGATTCCTTAATAATC [A/T] TTGGCTGGGGGTCTTTCTAA
85-12F4-86-2	TACTTCTTGAGGAAGCAGGT[G/C]AAAATTAACAAGAGCAATGG
85-12F4-L1A	TTCCTTAATAATCATTGGCT(G/T)GGGGTCTTTCTAACTATAAG
85-12F4-L1B	TTCCTTAATAATCATTGGCT[G/T]GGGGTCTTTCTAACTATAAA
L1-12G5-N1A	GCTTGAATAACGATGTCTAC [A/T] CTGCCTCGGCGTACGGCGGA
L1-12G5-N1B	GCTTGAATAACGATGTCTAC (A/T) CTGCCTCGGCGTACGGTGAT
85-12G6-86	AAAAGGGTACAANCTANTAA [T/G] TGATGACTCAACTTTCANTT
L3-12G8-L6	GCAAAGCTAAAAAATCGAC (A/G) AGTGTTCCTTACTACGCTCC

85-12G8-L1	CTAAAAAGATCGACGAGTGT [C/T] CCTTACTACGCTCCATCTAT
L5-12G8-L6	ACTCGGAGCAAAGCTAAAAA [G/A] ATCGACGAGTGTTCCTTACT
85-12G9-L3	GGTGGGTTTACCGTATCATT (T/C) GATCCATTGGATGGATCGAG
L6-12G9-N1-1A	AGGTGGGTTTAGCGTGGCAT (T/C) CGATCCATTGGATGGATCCA
L6-12G9-N1-1B	AAGTGGGTTTAGCGTGGCAT [T/C] CGATCCATTGGATGGATCCA
L6-12G9-N1-1C	AGGTGGGTTTAGCGTGGCAT (T/C) CGATCCATTGGATGGATCGA
L6-12G9-N1-1D	AAGTGGGTTTAGCGTGGCAT (T/C) CGATCCATTGGATGGATCGA
L6-12G9-N1-2A	TTCGATCCATTGGATGGATC(C/G)AGCATTGTGGATACAAACTT
L6-12G9-N1-2B	TCCGATCCATTGGATGGATC[C/G]AGCATTGTGGATACAAACTT
85/12G9/L3-1	GATCCATTGGATGGATCGAG (T/C) ATTGTGGATACAAACTTCAC
85/12G9/L3-2	TTGGGGTTTGGCCTGGTGAC(C/A)ATTTAACCGGACTCACGGGA
N1/13A3/N2A	CTGTCTCAGTTTGTTGGATC[C/G]AAATCGAAAGCGTAC
N1/13A3/N2B	CTGTCTCATTTTGTTGGATC[C/G]AAATCGAATCGAAAGCGTAC
L4-13A3-L5	ATGGAAAGTATGAATCTTTA (G/C)TCCACCCATTGTTCGCATTT
L4-13A3-N2A	TTTGTCCTGGTTCTGTCTCA (G/T) TTTGTTGGATCGAAATCGAA
L4-13A3-N2B	TTTGTCCTGGTTCTGTCTCA [G/T]TTTGTTGGATCGAAATCAAA
L4-13B6-N2-1A	TCTGAATAGGTCTTGGGGTT [A/T] TGTAAATTGTGTTGCGGGTG
L4-13B6-N2-1B	TCTGAATAGGTCTTGGGGTT [A/T] TGTAAATTGTGTTGCGGGTT
L4-13B6-N2-2A	ATGTAAATTGTGTTGCGGGT [G/T]TCTGAAAAGGGATTGGCGTT
L4-13B6-N2-2B	TTGTAAATTGTGTTGCGGGT (G/T)TCTGAAAAGGGATTGGCGTT
L3-13B6-L2	CTCTTCTTAGTCAATCTGAG (A/)AAGCCTGACGTCTCCTACAA
L2-13E8-N2	ACACTGTTGGAGGACGTGAA [T/G] AAGATATTCAAGACAACATC
N1/13F6/N2-1A	TCGAGGAACTGGAGATGGAT[A/G]AGGTAAACCTTTTGTTTTAT
N1/13F6/N2-1A	TCGAGGAACTGGATGGAT[A/G]AGGTAAACCTTTTGCTTTAT
N1/13F6/N2-2	TCTTTCGTATCTTGCTGAGT [C/T]GTTACGCCTGTCAACACCCG
L3-13F8-N2	GATCATAGTATCCGCCGGAA [G/C]CCTAGGGAGGCCACAGCTCC
L2-13F8-N2-1	GGAACCCTAGGGAGCCCACA (T/G) CTCCTTATGCTAAGCGGCGT
L2-13F8-N2-2	TAAGCGGCGTCGGGCCATCA (T/G)CTCAACTACAGGCCCAAAAT
85/1485/86	TTCCCCCCCCTCGATCCGGGC [A/G] GAAGACATTGTCAGGTGANN
86-14B5-L3	ACCCCTTCTTTTAGACCCAA (A/G) ACTCGCTTCGGCGGGTCGAT
N1/14C2/N2	GCACCAACATTGTAAACCTA [T/G]AGCTTCTTCCTCAGCCACCT
N1-14C2-L2A	AGCTTCTTCCTCAGCCACCT[T/G]CAACGAGAGCTCCTGGAAAC
N1-14C2-L2B	ATCTTCTTCCTCAGCCACCT[T/G]CAACGAGAGCTCCTGGAAAC
N2-14C2-85-1A	CCTCTATCCGCCATGGTTGC [A/T]CCAACATTGTGAACCTAGAG
N2-14C2-85-1B	CCTCTATCCGCCATGGTTGC (A/T)CCAACATTGTGAACCTAAAG
N2-14C2-85-1C	CATCTATCCGCCATGGTTGC (A/T) CCAACATTGTGAACCTAGAG
N2-14C2-85-1D	CATCTATCCGCCATGGTTGC (A/T) CCAACATTGTGAACCTAAAG
N2-14C2-85-2A	GCACCAACATTGTGAACCTA (G/A) AGCTTCTTCCTCAGCCACCT
N2-14C2-85-2B	GCTCCAACATTGTGAACCTA (G/A) AGCTTCTTCCTCAGCCACCT
N2-14C2-85-2A	CGAGAGCTCCTGGAAACCCT[T/G]GGCCACAGGTTTGTTTCTAT
N2-14C2-85-2A	GAAGAGCTCCTGGAAACCCT[T/G]GGCCACAGGTTTGTTTCTAT

N2-14C2-86A	AGCTTCTTCCTCAGCCACCT[T/AA]CAACGAGAGCTCCTGGAAAC
N2-14C2-86B	AGCTTCTTCCTCAGCCACCT[T/AA]CAACGAGAGCTCCTGGAACC
85/14C2/86A	AGTACATAGCTATTGACTAA [C/G] TTAAGTTCCTTGTATTGTTG
85/14C2/86B	ATTAAATAGCTATTGACTAA [C/G]TTAAGTTCCTTGTATTGTTG
85/14C2/86C	AGTAACTAGCTATTGACTAA (C/G) TTAAGTTCCTTGTATTGTTG
85/14C2/86D	ATTAACTAGCTATTGACTAA(C/G)TTAACTTCCTTGTATTGTTG
L1-14C2-85A	GAAGAGCTCCTGGAAACCCT[T/G]GGCCACAGGTTTGTTTCGGT
L1-14C2-85B	CGAGAGCTCCTGGAAACCCT (T/G) GGCCACAGGTTTGTTTCGGT
L1-14C2-85C	GAAGACCTCCTGGAAACCCT[T/G]GGCCACAGGTTTGTTTCTGG
L1-14C2-85D	CGAGAGCTCCTGGAAACCCT[T/G]GGCCACAGGTTTGTTTCTGG
85/14E2/86-1	AGGCTGCCCTCTCCCAATTC (A/C) AAAGCCAACTCCTAAACCAA
85/14E2/86-2	TTGACCCTCGGCAAGCCACC [G/T]GTCAAGCCATGCTGCAGCCT
85-14E8-86A	AAACATGGAAAGGCCTGATA (/G)TCACCGTCAAGCTCACCGTC
85-14E8-86B	AAAGATGGAAAGGCCTGATA [/G]TCACCGTCAAGCTCACCGTC
85/14E12/86	CAACCTGAAAAATTGTTTTA (C/A) CAACGGCCCCGCTTTCTCCA
85/14H10/86	TCACCGGCTTGAAGTCTTCC (G/T) CTGCATTCCCAGTCACCCGC
L1-14H10-86	AAGGCCAACAACGACATTAC (T/C) TCCATCGTTAGCAACGGAGG
B5/15A6/86	ACTCAGCTTTCTTATGCCTC [G/]ACTTGCGACACGCACTCCA
	·
85-15C4-86	TCCGCTAACATCTCTGCTG[G/T]TCACCTTAACCCAGCCGTAN
N1/15C10/N2A	ACGGTGAGATCTAACGGCGG [G/C] GATCCTTCAGTCCATAGTCG
N1/15C10/N2B	ACGCCGAGATCTAACGGCGG [G/C] GATCCTTCAGTCCATAGTCG
N1/15C10/N2C	TCGGTGAGATCTAACGGCGG [G/C] GATCCTTCAGTCCATAGTCG
N1/15C10/N2D	TCGGCGAGATCTAACGCCGG[G/C]GATCCTTCAGTCCATAGTCG
Warascaora	
86-15E5-N2	CCCGCGAGGAGCACGACTAC (A/T) GATTCTCCGTTTTCAAATCC
N1/15E5/N2-1	CTGTTCAAGAAGAAGTTCGG[C/T]AAGGTCTACGCTTCCCGCGA
N1/15E5/N2-2	CCCTCTGCTCGTCACGGCGT [T/A] ACGCAGTTCTCGGATCTGAC
85/15E5/86-1	CGAGGATCACTTCTCTGT (G/T) CAAGAAGAAGTTCGGCAAGG
85/15E5/86-2	GAAGAAGTTCGGCAAGGTCT[GA/AC]GCTTCCCGCGAGGAGCACGA
L1-15E9-86A	TCCACTCGCCGGGAAGAAC(A/T)CGACAAACCGTTGTCTACTT
N1-15E9-86A	ATCTTCCTCTCTATCTTCAA(C/T)GTCGTGACAAGAATGATGTG
N1-15E9-86B	ATCTGCCTCTCTATCTTCAA (C/T) GTCGTGACAAGAATGATGTG
N1-15E9-66C	ATCTTCCTCTATCTCAA (C/T) GTCGTGACAAGAATGAGTGG
N1-15E9-86D	ATCTGCCTCTCTATCTTCAA (C/T) GTCGTGACAAGAATGAGTGG
L1-15E9-86B	TCCACTCGCCGGGAAGAAC (A/T) CGACAAACCGTTGTCCACTT
L1-15E9-86C	TCCACTCGCCGGGAAGAAC(A/T)CGACAAACCGTTGTCTACGT
L1-15E9-86D	TCCACTCGCCGGGAAGAAC(A/T)CGACAAACCGTTGTCCACGT
N2-15E9-85A	TCCACTCGCCGGGAAGAAC(A/C)CGACAAACCGTTGTCTACTT
N2-15E9-85B	TCCACTCGCCGGGAAGAAC(A/C)CGACAAACCGTTGTCTACAT
N2-15E9-L1	ATGGCTCGCGACGGGTCTCC [G/T] GTAAACCTCGGAGAGCAGAT
N2-15E9-86	GCCGACTCTCGAAGCTTCTT(A/)ACTCCACTCGCCGGGAAGAA

85/1529/86-1	GAATCTAGGAGAGCAGATCT[T/G]CCTCTCTATCTTCAATGTTC
85/15E9/86-2	TCCACTCGCCGGGAAGAAC(C/T)CGACAAACCGTTGTCTACAT
85/15E9/86-2B	TCCACTCGCCGGGAAGAAC(C/T)CGACAAACCGTTGTCCACAT
85/15E9/86-2C	TCCACTCGCCGGAAGAAAC [C/T] CGACAAACCGTTGTCCACGT
85/15E9/86-2D	TCCACTCGCCGGGAAGAAAC [C/T] CGACAAACCGTTGTCTACGT
85/15F1/86	GCAGGTAAAATTCTACAGAC (C/A)TTCCCTTTTCATTGTAGTTA
85/15F5/86	TCTCCTCCGCCGCGCAAGAA [G/A] AAATCGACAGCGGCGCGTCT
85/15F10/86A	GTGCCCTAAAGATACCCTCA (A/G) GCTTGGTGTCTGCGCTAATG
85/15F10/86B	TTGCCCTAAAGATACCCTCA (A/G) GCTTGGTGTCTGCGCTAATG
N1-15E9-N2-1	GTCATGAAGATATTCACTAC (A/G) CCGACTCTCGAAGCTTCTTA
N1-15E9-N2-2	GCCGACTCTCGAAGCTTCTT[/A]ACTCCACTCGCCGGGAAGAA
£/	
N2-15G1-L3	TTCTTCCCACAGGTGAAACT (T/C)GCTAACTTCCTTCCAAAGTA
86-15H7-N2	CAGGACAATGTGTTTGTGAC (A/T) GTGGTTGCATCCATTCAATA
N1/15H7/N2	TATGTATCAGGACAATGTGT (GA/TT)GTGACTGTGGTTGCATCCAT
N1/15C10/N2	ACGGTGAGATCTAACGGCGG[G/C]GATCCTTCAGTCCATAGTCG
N1/16D6/N2	GCTAAGGTAGTTGGAGGAGC [CAA / GTG] CCACAGCCACGCGACTAAGG
11710007112	0011110011101100110011011011011010101010
N1/16A1/N2-1	GCTAAGCTACGCAACTGCCA(C/T)CAATCAGGGCAAGCTAAAGG
N1/16A1/N2-2	ATCAGGGCAAGCTAAAGGAA(C/T)GAATGACATTGAAGATGTGA
111/10/11/112-2	The district of the last of th
85/16A5/86	TATACACTCTTTAAAAGCGT [G/C] TGTGTGTACCCATCTCTCTT
83/1083/00	
85/16B6/N2A	GACATTTCTAGACTTGAGA [T/] TGGCTGCGTATTGGCTGTCT
85/16B6/N2B	GGCATTTCTAGACTTGAGA [T/] TGGCTGCGTATTGGCTGTCT
85/16B6/N2C	GGACATTTCTAGACTTGAGA (T/)TGGCTGCGTATTGGCTGTCT
85/16B6/N2D	GGGCATTTCTAGACTTGAGA (T/) TGGCTGCGTATTGGCTGTCT
85/16B6/N1	GGATCCATCTCAACTATGGT [A/C] GTATTATCGTTGAGGCTAGG
85/16B6/N1	TGGACCATCTCAACTATGGT [A/C]GTATTATCGTTGAGGCTAGG
N1/16B6/N2	ATGGCTGCGTATTGGCTGTC [C/T] AAGGCTGGATCTTGGTCCCA
11710207112	NAGOCIOCANTICOCIONA (O'N) INGGOCIOCANTICOCIO
05/1607/06	GTATGTGATTCGGAAGAGAA [T/]CAAACTAAGTGCCGAGAAAG
85/16B7/86	ATMIGITALITY CONTRIBUTION (11) CARROLLING CONTRIBUTION
96 / 1 6D6 / NO	GCTAAGGTAGTTGGAGGAGC [CAA/GTG] CCACAGCCACGCGACTAAGG
86/16D6/N2	GA SANGO TAO TAO NAMBO LANGO TANGO
95/16D10/06	CTCAACGTAGCAAGTAATAA [T/G]ATACTGTCTATTTATGGTTA
85/16D10/86	CTCAACGTAGCAAGTAATAATTAGTAATTACGTTA
N1 /1 CEO (N2)	1010mmmcccc1mmcmmmcrm/11ccc1
N1/16E9/N2A	AGACTTTCCCCATTCTCTTC[T/A]CCATCCACCGTCGAAACCCA
N1/16E9/N2B	TCACTTTCCCCATTCTCTTC [T/A] CCATCCACCGTCGAAACCCA
05 /1 502 /05 41	10000011100001110001111010011111101001111
85/16H3/85-1A	ACTTCGAAACTGTAAACCTA [A/T] ACTTTAAGAGTTTAGAGCTA
85/16H3/86-1B	ATTTCGAAACTGTAAACCTA [A/T] ACTTTAAGAGTTTAGAGCTA

B5/16H3/86-2a	CACCATCGGAGAAAGAGGTA[C/T]TTCGAAACTGTAAACCTAAA
85/16H3/86-2B	CACCATCGGAGAAAGAGGTA [C/T]TTCGAAACTGTAAACCTATA
85/17A5/86	CTAAGGCGTCTCCTGAAGAA [A/G]TACAGAGAGTCGAAGAAGAT
85/17C7/86A	CCGCGGACGACGCTTTCTTC(C/A)TCTGCTCCACCGCGAGCGCC
85/17C7/86B	GGGCGGACGACGCTTTCTTC [C/A]TCTGCTCCACCGCGAGCGCC
85/17F7/86	QAGGAGTAGTCTCCATGGCC (G/) AAGAAGAGCGTCGGAGACCTG
85/17G12/86 .	GAAGTTAGGGCTTCTAAGAT[C/T]AAGTTCGGCAAGGCTTTAAC
85/182/86	TCAAAACTAATATTTCTTTT[G/C]TTGATTGGTAATAAACAGGT
85/18A11/86	TTCCAGTGAAAAGGCATTGT (T/G) CTCCAAAATCTCGCTCTGCG
85/18A11/86	TCTGCTGAGACTGTGGCACC[/G]ATCTCGTTCTACGCCGGCTTC
85/18F5/86	AAGCAGCTCTGACTTGAATG (C/A) GAGAGGTTAATCAGACTGTG
85/18F5/86	AACTTGTTCTCTGATGGTGT(A/T)GTAAGTACGTGCGCCCCTTT
85/18H10/86-1	GTTTTTCTTTTGTTTT [A/C]TTAACTAGCGACTTGAAACT
85/18H10/86-2A	CATATTTTTCTCTCCCTTA [G/A] ATTGAAGCAATCAAGAAGAT
85/18H10/86-2B	CATATTTTTCTCTCCCTTA(G/A)ATTGAAGCAATCAAGAAAAT
85/18H10/86-3A	TAGATTGAAGCAATCAAGAA[G/A]ATCTCAGACTTCATCACCCA
85/18H10/86-3B	TARATTGAAGCAATCAAGAA [G/A] ATCTCAGACTTCATCACCCA
25.44.227.426	GCATCCAACTCCAAGGATGA (/C) CCTGCCAAGGTGCTGCTAACT
85/19B3/86	CATCOMC COMMON ON (7 C) COT GOOD AND COT GOO
85/1908/86	QAGCTCAGGGATGGTGGATC (A/T) GACTACCTTGGAAAGGGTGT
8371308700	
N1/19F4/N2	TGGGGTTAGTCGAAATAGGT(A/T)AAATGCTTTGAGTATGTGTA
N1/19H1/N2	TACGCGCAGCACGGACTTGC G/A ACGCAAGCAATCGAGCTTTT
95 /2094 /95 -1	GAAGCCCATGGTACGGAGCG[G/A]GAGAGAGTCAAGTACTTGGG
85/20B4/86-1	GAAAACGTCGCCAAGCCGAA[/G]GGGTCCATCAGGAAGCCCATG
85/2084/86-2A 85/2084/86-2B	GTGAACGTCGCCAAGCCGAA[/G]GGGTCCATCAGGAAGCCCATG
63/2084/80-28	OTOMICS COCCINIOCES IN () O O O O O O O O O O O O O O O O O O
N1/20B12/N2	AACGGGTCACTGCTAAATCA [T/A] AAGGATCACAAGGCTGGGAC
85/20C12/86	CTAGCCTACTTTGGGAAAAG[/T]TTCGTTATTGTTTTGTGTGG
85/20D2/86	GACTTCAAGGACTTCGCCGG [A/C] AAATGCTCCGACGCTGTCAA
N1/20D2/86	CGCGGGAGTGCTCACCTCCG (T/G) TCGACTCAGACCGTCGGCAA
85/20D3-D6/86-1A	AAGAGGCTGAGGGGCTAAA (C/G]AGCAGGAAGAGCCTGAAGGA
85/20D3-D6/86-1B	GAGAGGCTGAGGGGCTAAA [C/G] AGCAGGAAGAGCCTGAAGGA
85/20D3-D6/86-1C	AAGAGGCTGAGGGGGCTAAA [C/G]AGCAGGAAGAGCCTGAAGGA
85/20D3-D6/86-1D	GAGAGGCTGAGGGGCTAAA (C/G) AGCAGGAAGAGCCTGAAGGA

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85/20D3/86-2A	GAGGAGGCTACATGCAGCT [G/A] AAGAGGCTGAGGGGGCTAAA
85/20D3/86-2B	AGGAGGGGTACATGCAGCT [G/A] AAGAGGCTGAGGGGGCTAAA
85/20D6/86-1	GGAGGGGCTACATGCAGCTA [G/A] AGAGGCTGAGGGGGCTAAAG
85/20D6/86-2	AGGGGGCTAAAGAGCAGGAA [G/C] AGCCTGAAGGAGAGTTGGAA
85/20D6/86-3A	GAGTTGGAACTGAATGATGT (T/C) CAACCTATGAAGAAGAAACA
85/20D€/86-3B	GAGTTGGAACTGAATGATGT (T/C) CAACCTATGAACAAGAAACA
85/20D6/86-4A	GATGTTCAACCTATGAAGAA (G/C) AAACACCGAGGACCAACGAG
85/20D6/86-4B	GATGTCCAACCTATGAAGAA [G/C] AAACACCGAGGACCAACGAA
85/20D6/86-5	AGATGAAACACATCGCCAAA[/G]GATCCAAACACTCGAGAAAGA
85/20D6/86-5	CCATTAGTGAGGGAGCATGT[T/A]CCTGTCACATTTGATGATTG
85/20D6/86-7	AGCAGGAAGAGCCTGAAGGA[C/G]AGTTGGAACTGAATGATGTT
85/20D6/86-8	AAACACATCGCCAAAGATCC[CG/AA]ACACTCGAGAAAGAGTCGAG
85/20D6/86-9	ATCCTGTTGGTGAAGGATCA (C/G) TGAATCTGTCTTCTTACTTG
N1/20D8/N2	CTCATAGGCGATCTGGAGTA [T/G] GCAAATCGAATCTCCTCTCC
85/20C12/86	CTAGCCTACTTTGGGAAAAG [/T] TTCGTTATTGTTTGTGTGG
N1/20E1/N2A	TGCACGCCTCACTTGTTCCT [T/A] CCAATCTGACATCAAGGATT
N1/20E1/N2B	TGCACGCCTCACTTGTTCCT[T/A]CCAATCTGACATCAAAGATT
N1/20F1/N2-1	NGTGTTTTTGAGGTGAAAGC [A/T] ACAAATGGAGATACCTTTTT
N1/20F1/N2-2	GANAAGACTTCGACAACACT[T/C]TGTTGGAGTNTTTGGTNTCT
	- Constitution of the cons
N1/BoC-a2/N3-1	CCCATCTCATCCTTTCTTGA [A/G] CCGTTGAATCAAGCTCCTGG
N1/BoC-a2/N3-2	CCCGAGCCATTAGGACAAGA(T/C)GACTTGCCGTTTGACCAAAC
N1/BoC-a2/N3-3	TACATTCTCATTGGTTGGTT(C/A)TTGGGAAATAAAGTACCAAC
85-SC3-L2	GCACGCGCTAGAGTTGTTGC [C/A] AGAAGGAATGAACAATCTGA
86-SC3-L1	ATGGACGTTAAAAACGCTTT(T/C)CTTCCCGGAAATCTTGCTGA
86-SC3-N4-1A	TCCCGGAAATCTTGCTGAAA(A/C)CGTTTACCTGCGACAACCAG
86-SC3-N4-1B	TCACGGAAATCTTGCTGAAA[A/C]CGTTTACCTGCGACAACCAG
L1-SC3-N3-1	ATCTTGCTGAAACCGTTTAC [C/A] TGCGACAACCAGCCGGTTTT
L3-SC3-NS-1	AACAAGCCTGATCACGTTTG (C/G) TTGCTTCACAAATCTTTGTA
N1-SC3-N3-1	GAAGGCGTCCACTATCTTGA (A/G) ACCTATAGTCCTGTTGTTCG
N1-SC3-N3-1	
N3-SC3-N4-1	GAAGGCGTCCACTATCTTGA (A/G) ACCTATAGTCCTGTAGTTCG CTTGAGACCTATAGTCCTGT (A/T) GTTCGGTCCGCCACAGTTCG
N3/SC3/N5-1	CACAGTTCGTACAGTTCTTC (A/C) CATTGCCACTGTTATGCACT
L2/SC3/N3-1	AACCAGCCGGTTTTGTTGAC(C/A)GAAACAAGCCTGATCACGTT
	TOTAL CONTROL TO A TOTAL CONTROL OF THE CONTROL OF
L3/BL1/N5-1	ATGTCTTCAAAGTGCTCTGT[C/T]GCAACGCACGTCCGAACAAG
16/BL1/N5-1	GTGCTCTGTTGCAACGCACG(C/T)CCGAACGAAGAAAGACAACGA
2A7/287-1	GACGAGATGGA AAGCAACCA (M/OLAGOGTTA) GAAGAAAA
2A7/2B7-2A	GACGAGATGGAAAGCAACCA[T/G]AGGGTTCAGGAGAACCGGCT
2A7/2B7-2B	GARARATGCARRATACTTAC [A/G] ACCTTGCTCARCARGCTART
2A7/2B7-3	GAAGAATGCAAAATACTTAC (A/G) ACCTTGCTCAACAAGCTAAT
	CCTATAAAAGGGCCACGAGT[G/T]GAGATTGGGGGTCGTGGGGT
2A7/2B7-3	TGATGATGGATCTCAGGCTA [T/G]TCCGCATGGTAAAAACTACA

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		7
85 86		1
2B7/2E7-1A	AGAGCACGCATTGGACCTAT(A/T)AAAGGGCCACGAGTTGAGAT	
2B7/2E7-1B	AGAGCACGCATTGGACCTAT (A/T) AAAGGGCCACGAGTGGAGAT	
2B7/2F7-1A	GCGGTTGCTGCAGCTCA (A/G) AQAGCACGCATTGGACCTAT	1
2B7/2F7-1B	TCGGTTGCTGCAGCTCA (A/G) AGAGCACGCATTGGACCTAT	
2B7/2F7-2A	ATGCAGGGCAAGAGAGGGAC (G/T) CGGTTGCTGCAGCTCAA]
2B7/2F7-2A	ATGCAGGGCAAGAGAGGGAC [G/T] CGCTTGCTGGAGCTCAG	
S71G2/G6-1	ATAATACTTGATATGCCATT[G/T]TGTCCTCTTATTTTTAACAT	ď
		ŀ
S67G1/G5-1	ATGGCCTCGTCGGCCACTGC (A/C) OTCGCTCCGTTCCATGGGCT]
S67G1/G3-1	GCCGCTCCTCCAGAAGCCTC (G/A) GCAACGTCAGCAACGGCGGA]
S67G1/G3-2	GTGTTGCCCATCCCA(A/T)TTCCCAACCCCAAACGAACC]
S67G1/G3-3	GTACCTGCCGCCGCTCTCGA [CG/AC] GGACGACCTCCTGAAGCACC	1
		1
S66G2/G3-1	AGTGAGCCCCCTTCTTATTC[/T]TAAGGTGATAGGTTTCTAAA]
S66G1/G3-1	AATGTAATGGTACTCCGCGC [T/C] ATGGCTCTCGTACTTAGGAA	1
S66G1/G2-1	AAATAGGCTCGGGCAATTAT [C/]CAGCTTAGGGACAGCAAGCG	1
		1
565G3/G6-1	TEEGCCCTGCCTCCGGTTTT(A/T)GCCCGACCTTCGAAACATTC	1
565G3/G6-2	ACCACTGACGTAGCACCTCC(G/T)ACTTCTCGTTGTAAAACCCC	1
S65G3/G5-1	GGAGGTTCGCCTCATCTTAT(C/T)GTTGACGAGCCACATCCACT	1
S65G4/G6-1	GCTCCGACTTCCAATCTTGA (A/C) CCTCCACCCTGCCTCCGGTT	1
		1
554G1/g3-1	CTGGTTGAAATGTGTTGAAG(C/A)TACTAGTGATGAACTGCTTG	1
S64G1/g3-2A	GCTGCTCCAAGCGAGCCCGC [C / G] CCGAAAAAGGAAAAAGGTGA	1
564G1/g3-2B	GCTGCTCCAAGCGAGCCCGC [C/G]CCGAAAAAGGAAAAGTTGA	1
S64G1/g3-3A	CGCCCCGAAAAAGGAAAAAG (G/T) TGAAGGTCCTTACTCACCGA	†
S64G1/g3-3E	CCCCCCAAAAAGGAAAAAG(C/T)TGAAGGTCCTTACTCACCGA	1
S64G1/a3-4A	GAACCGGCCACAGTGCCTGA[T/A]TTTGGCGGTGAGACCTCTTC	1
864G1/g3-4A	GAACCGCCCACAGTGCCTGA(T/A)TTTGGCGGTGAGACTTCTTC	1
B04017 43-4X	- ONNEGOCCACAGO I GEORGIA (1/A) 1110 GCGG I GAGACII CITC	1
S63G5/G6-1	CARTTOTTACCTGAGCAAGA [T/]TTTTGTGTACTTGACTTGTT	┨
S63G4/G6-1	TACTGAGAGAATGCAACATC [C/G] AGCATTCTGTGATTGGAGTC	1
S63G4/G5-1A	TTTTAGTGTACTTGACTTGT [C/T] CTCCTCCACAGATGAAATAT	1
		┨
S63G4/G5-1E	TTTTTGTGTACTTGACTTGT[C/T]CTCCTCCACAGATGAAATAT	1
S63G3/G6-1	TCTGTGATTGGAGTCTGCTC[G/A]CGTGTCAGCTCTGGATGTGA	·
CERCE ICE		1
S57G5/G6-1	AACTACAAAAAGCATCTCCT G/T GGATTTGGCTATCTCCTTTT	ŀ
S57G2/G5-1	TTAGCGCGAAAAAAAACTC[/T]TTTTTTTTTTTTTTACT	ŀ
S57G2/G3-1	ITCAATCCAATCAATTTAATT [T/C] CTTCCTTTAAAAAATATTATC	1
S57G1/G2-1	TTACTACGAAAACTCTTGA[G/T]TCTAGGAATTTGAATTTGTG	1
S57G1/G2-2A	CTTCTTGGATTTTGCTATCT T/C CTTTTACTACGAAAAACTCT	1
S57G1/G2-2B	CTCCTTGGATTTTGCTATCT [T/C] CTTTTACTACGAAAAACTCT	1
S57G1/G2-3A	TTTTACTACGAAAAGCATCT T/C;CTTGGATTTTGCTATCTTCT	ł
S57G1/G2-32	ITTTACTACGAAAAGCATCT [T/C] CTTGGATTTTGCTATCTCCT	1
557G1/G2-4	GAAGCCAAATCCTATTATTT (T/C) CTGCCTCTAGGGTCTGAATG	4
		-
IS56G4/G6-1	IGTACACTGTTACAATCACAC [T/G]TAGTGAAGCGCAACACAGAT	į

- Marker (Stout)

S56G4/G6-2	GCCTTATCATCCTCTAGGTA (T/A) TGGAGACGAGTGACCAGTCT
S56G4/G6-3	CTTTTCTTCAGACCCGAGCC [C/T] CCAATCGCGCCCTTCTGTGC
S56G4/G6-3	CTTTTCTTCAGACCCGAGCC (C/T) CCAATCGCGCCCTTTTGTGC
556G4/G5-1A	GAGCCCCAATCGCGCCCTT (C/T) TGTGCCTTGGCCTTGAGCTC
S56G4/G5-1A	GAGCCTCCAATCGCGCCCTT(C/T)TGTGCCTTGGCCTTGAGCTC
S55G1/G3-1	GAAGGAGCAGCGCAAGG (A/) ACGTGTTCCAAGTCAACGTC
S54G2/G3-1	GTAGAAAGTTAGCAAAAACA (T/) TTTTTTAGTGAAAAAACATA
S54G2/G3-2	ATTGTGGCTAGAAACTTTGG[/T]TTTTTTTAAATTATGGTCAT
S53G5/G6-1	GCAAACCAACACCAATCTTC[G/C]AAATGAGCAAAGCAGAGACT
S53G5/G6-2	CAGATCGGTTGTCCTCAGAG[A/]AAGTCACCTACCTGCAAACC
S53G5/G6-3	AATTCTACATAGGAGTCATG[C/T]ACAAGTACTTGTTTAAAGGA
S\$3G5/G6-4	ACAAGTACTTGTTTAAAGGA [C/]CATGCCGGAATACACGCTGC
S53G5/G6-5A	GAGCGAGATCGATCCTGTTG[T/C]CATCCATCACTGCCATAGGA
S53G5/G6-5B	GAGCGAGATCGATCCTGTTG (T/C) CATCCATCACTGCCGTAGGA
S53G4/G6-1	TAGTCATAGCAACAGCATGC [G/A] TCGTGATGTAGCGTTCACCC
S53G4/G6-2	CANTTGANGAGGANANANA [/T]TCTACATAGGAGTCATGTAC
S53G4/G5-1	CAGAGACTCCACAAGGCGAA [A/C] GGAGTCCACAATAGTTCGTC
S53G3/G5-1	CCCACGGCGGAGATGGTGG [T/]TAGAAGCGGAACCACCGAGC
553G2/G6-1	ACTTGTTTAAAGGACATGCC[G/]GGAATACACGCTGCCCAGGC
S53G2/G3-1	CCCAGGCCTTCCCACGGCGG [A/G]GATGGTGGTTAGAAGCGGAA
S53G1/G6-1	CAAAGCAGAGACTCCACAAG (A/G)CGAACAGAGTCCGCAATAGT
S53G1/G6-2	GAACAGAGTCCGCAATAGTT (T/C) ATCCTAATGCTACTTCGAGC
S49G2/G6-1	TGCTGCTGCACTTGCTCATC [G/C]TTACTGATTTGCTGAAATGT
S49G2/G4-3	TGCTGTAAAAAACGAGTGGA [C/G] TCCTAGTGTGTCGTGCGCTG
\$49Q2/G4-3B	CGCTGTAAAAAACGAGTGGA [C/G] TCCTAGTGTGTCGTGCGCTG
S49G1/G5-1	TGGCTCTAGCGAAGCGTAAA [GA/]GAGCAACGAAAGCAAGCGAG
94906/01-3	TTTGCGCGCAATAAATCAGA (A/G) AGCTGATCTGAATCTGACCC
S49G5/G1-2	CAAGCAAGCAAGCTGTCTGT [C/T] CGTATGTGTCTGGCATGTTA
S49G1/G4-1	CGCTGTAAAAACGAGTGGA [C/G] TCCTAGTGTGTGCGCGCTG
S49G1/G4-2	TCTAGCGAAGCGTAAAGAGA [G/T] CAACGAAAGCAAGCGAGCGA
S49G1/G4-2B	TCTAGCGAAGCGTAAAGAGA [G/T] CAACGAAAGCAAGCGAGCAA
S49G4/G1-1	GTAAAGAGATCAACGAAAGC [A/G] AGCGAGCAAGCAAGCTGTCT
S49G1/G3-3	TGATTTGCTGAAATGTTGGC [G/T]TTTTTTTTTGGTTGTTGGCCG
S49G1/G3-3B	TGATTTGCTGAAATGTTGGC [G/T]TTTTTCTTGGTTGTTGCCGG
S49G4/G6-2	CGCTGTAAAAAACGAGTGGA (G/C) TCCTAGTGTGTCGTGCGCTG
S49G4/G6-2B	TGCTGTAAAAACGAGTGGA (G/C) TCCTAGTGTGTCGTGCGCTG
S49G4/G6-3	CTCTAGCGAAGCGTAAAGAG[AT/]CAACGAAAGCAAGCGAGCGA
549G4/G6-3B	CTCTAGCGAAGCGTAAAGAG(AT/)CAACGAAAGCAAGCGAGCAA
S49G4/G5-1	CGGTGACCCGATGATTATGG [T/C] GGCGGCCACACCTGCAATGA
S49G4/G5-1B	CGGTGACCCGATGATTATGG [T/C] GGCGGCCACACCTCAAATGA
S49G4/G2-1	CGCTGTAAAAACGAGTGGA [G/C] TCCTAGTGTGTCGTGCGCTG
549G1/G6-1	TTTGCGCGCAATAAATCAGA (G/A) AGCTGATCTGAATCTGACCC
S49G1/G6-2	TGCTGCTGCACTTGCTCATC [G/C] TTACTGATTTGCTGAAATGT
S49G1/G6-3	TGGCTCTAGCGAAGCGTAAA [GA/] GAGCAACGAAAGCAAGCGAG
S49G1/G5-1	TGATTTGCTGAAATGTTGGC [G/]TTTTTCTTGGTTGTTGGCCG

S49G1/G4-1	TCTAGCGAAGCGTAAAGAGA [G/T] CAACGAAAGCAAGCGAGCGA
549G1/G4-1	TCTAGCGAAGCGTAAAGAGA [G/T] CAACGAAAGCAAGCGAGCAA
S49G3/G6~1	CCACCCGTTGCAGTGCTGT[/T]GCTGCTGCACTTGCTCATCG
S48G3/G6-1	GATTCAGAAACAGTGGCGGC (A/G)GATGTAGCATCAACACGCCC
S47G1/G3-1A	CACTCCTCCAAGATCTTATG[/A]AATCGTTCTGAGCTTATCGG
547G1/G3-1B	CACTCCTCCAAGATCTTATG[/A]AATCCTTCTGAGCTTATCGC
S47G1/G3-2	GCAAGCTGGAGCACGGTACT (A/G) TAGTAGCGGCCGGCGAGGGA
S47G1/G3-3	GGAATACTCGATAGGCTCCC (A/G)CTGTGGGTAACAGTATTCCT
S47G1/G3-4	CTCCTCGTGGTAGTGACGAT [G/C] ATTGCATCGGTGCCACAGGC
346G5/G6-1	CCCCCACCAGCACCACCGG[/CC]CCCCGGCCAATTGGACCCAA
946G3/G6-1A	GAAACAAGGGATAAAATGGG[G/A]AAAAAATCATTTTCCCTGAC
S46G3/G6-1B	GAAACAAGGGATAAAATGGG[G/A]AAAAAATCATTTTCCCTAAC
S46G3/G6-2A	GGGAAAAATCATTTCCCT(G/A)ACCTTTACCACCACCATTAC
S46G3/G6-2B	GGAAAAAATCATTTTCCCT(G/A)ACCTTTACCACCACCATTAC
S46G3/G6-3	CAATTGGACCCAAACCAAAA (A/C) CAAACCCAACGGCTTTTCCT
S46G2/G6-1	CGCGGTTTTCCCGGCGGCTA (T/C) GCGGGCGGGCCGCCGGCCCA
S46G1/G6-1A	GCGGGCGGGCCGGCCCA (A/T) CGTAACCTACCACCGTATTT
S46G1/G6-1B	GCGGGCGGCCGCCCCA(A/T)CGTAACCTACCACCGTACTT
S46G1/G2-1	TTCGGTGTCACTGACCTGTA [G/A] CATCAGCAGTAGCAGCGCCC
34001702-1	
S45G5/G6-1	ATGAGTATATTCAAGTCATA[T/C]TGTGAACTAGAATGTTATTT
S45G5/G6-2A	CCTAGACGCTGACCGCCACA [G/A] ACGGCGGCGGCTGCCAAATC
S45G5/G6-2B	CCTARACGCTGACCGCCACA [G/A] ACGGCGGCGGCTGCCARATC
545G5/G6-3	TGAACAAACCATGCGCTACC[C/T]AGCTAGGTGTTTTAAAGTAA
S45G4/G6-1	TCCGCGGAAACAACATCCGA (G/T)TTCTTGAGGATAACCCAGCT
S45G4/G5-1	GGGAGGGGAAAAAAAAGAA [G/A] AGCGTTGGTTGCGGTTCAGT
\$45G4/G5-2	GGCGGCTGCCAAATCCGCGG[/A]AAACGACATCCGAGTTCTTG
S45G2/G4-1A	CTAGAATGTTATTTCTTCAC [C/A]GTTGACCATGGAAAAAAACA
S45G2/G4-1B	CTAGAATGTTATTTCTTCAC [C/A] GTTGACCATGGAAAGAAACA
S45G2/G4-2A	TTCACCGTTGACCATGGAAA [A/G] AAACAGTAATAAGTTCTTGT
S45G2/G4-2B	TTCACAGTTGACCATGGAAA (A/G) AAACAGTAATAAGTTCTTGT
S45G1/G6-1	TTCTTCACAGTTGACCATGG[/A]AAAAAACAGTAATAAGTTC
545G1/G5-1A	GAACCCACCGTGCCCTGGGA[/G]GGGAAAAAAAAAAAAGAGAGCG
	GAACCCACCGTGCCCTGGGA [/G]GGGAAAAAAAAAAAAAAAGCG
S45G1/G5-1B S45G1/G5-2A	TGGGAGGGAAAAAAAGAA [G/A] AGCGTTGGTTGCGGTTCAGT
	CGTACCAGCTAGGAATCGTA[A/G]AAAAGCCTAGACGCTGACCG
S45G1/G5-3	COMMONIA
S44G5/G6-1	GCTGCGTCAATCACTTC[T/A]CCCACAGGCGTCAAGTACAG
544G3/G4-1	GACAGATTCCAAAGTAGTCG[C/T]CGGCCAGGTCGAAAAAGAAT
S44G2/G6-1	GGCGCTGCGTCAATCATCAC(A/T)TCACCCACAGGCGTCAAGTA
S44G2/G4-1A	TCGGTGTCACCACATGCATA (T/C)TCAGGACAGATTCCAAACTA
S44G2/G4-1B	TCGGTGTCACCACATGCATA [T/G]TCAGGACAGATTCCAAAGTA
S44G2/G4-2A	GTCGCCGGCCAGGTCGAAAA(G/A)GAATACTCAGCAAAAGACCC
S44G2/G4-2B	GTCGTCGGCCAGGTCGAAAA [G/A]GAATACTCAGCAAAAGACCC
S44G2/G3-1A	TATTCAGGACAGATTCCAAA [C/G]TAGTCGCCGGCCAGGTCGAA
S44G2/G3-1B	TAGTCAGGACAGATTCCAAA C/G TAGTCGCCGGCCAGGTCGAA

S44G1/G6-1	GCGTCAAGTACAGATACGCA(A/G)CACGCCTCAGCTTCGCCTTG
S44G1/G2-1	CCTGGGACTCCGCAAATTGC (G/A) AGCACTCGGTGTCACCACAT
S43G2/G6-1	GCTGGTTCATTATCTGACCT [G/T] GATTGCATTGCAGCTACAAG
543G2/G6-2	CTGGATTGCATTGCAGCTAC (A/G)AGAAGCCCGTGGAAGGCCGG
S43G2/G6-1B	GCTGGTTCATTATCTGACCT[G/T]GATTGCATTGCAGCTACGAG
S43G2/G6-2B	CTTGATTGCATTGCAGCTAC [A/G] AGAAGCCCGTGGAAGGCCGG
S43G2/G6-3	TCAGCCCCTACTACGCCGAA[G/]GAGCTCATCTCCGGCATCGC
S43G2/G5-1	TACCCGGAGCTGAACCTCCC[C/G]GAGAGATTCAAGTCCTCCTT
543G2/G4-1	TGCATGTGAACATTCATGAA[T/C]GGTAACCCACAACTGTTCGC
543G6/G1-1	CTCCTACCAGGGCCGGTTCG[T/]CCTTCTCCGACTACCCGGAG
S43G1/G6-1	TGAATGGTAACCCACAACTG(C/T)TCGCGTCCTGCTGGTTCATT
S43G1/G5-1	GCCGACAGGGTCCTCACCGT[G/C]AGCCCCTACTACGCCGAAGA
S43G2/G6-1	GCTGGTTCATTATCTGACCT[G/T]GATTGCATTGCAGCTACAAG
543G2/G6-1B	GCTGGTTCATTATCTGACCT[G/T]GATTGCATTGCAGCTACGAG
543G2/G6-2	CTGGATTGCATTGCAGCTAC [A/G] AGAAGCCCGTGGAAGGCCGG
S43G2/G6-2B	CTTGATTGCATTGCAGCTAC (A/G) AGAAGCCCGTGGAAGGCCGG
S43G2/G6-5	TCAGCCCCTACTACGCCGAA [G/]GAGCTCATCTCCGGCATCGC
S43G2/G6-3	TGCATGTGAACATTCATGAA (T/C) GGTAACCCACAACTGTTCGC
	CTGGTGGTGGTCCTTCTCTG[AAAC/]TGAAACTGAAACTGACTGCA
\$43G2/G3-1	GACCATCTTCACGTACTACC(TACC/)AGACCGCTTTCTGCATCCAC
543G2/G3-3	CTGACCATCTTCACGTACTA (CCTA/) CCAGACCGCTTCTGCATCC
543G1/G6-1	
543G6/G1-1	CTCCTACCAGGCCGGTTCG (T/)CCTTCTCCGACTACCCGGAG
S43G6/G1-1	GAGATTCAAGTCGTCCTTCG(G/)ATTTCATCGACGGGTCTGTT
S43G1/G6-2	TGAATGGTAACCCACAACTG [C/T] TCGCGTCCTGCTGCTTCATT
543G1/G6-3	GCCGACAGGGTCCTCACCGT[G/C]AGCCCCTACTACGCCGAAGA
S43G1/G5-1	TCTGACCATCTTCACGTACT [ACCT/]ACCAGACCGCTTTCTGCATC
543G1/G4-1	CTTGATTGCAGTAC [G/A] AGAAGCCCGTGGAAGGCCGG
S43G1/G4-1	CTGGATTGCATTGCAGCTAC [G/A] AGAAGCCCGTGGAAGGCCGG
543G1/G3-1	GCTGGTTCATTATCTGACCT [T/G] GATTGCATTGCAGCTACGAG
S43G5/G6-1	AGAGATTCAAGTCGTCCTTC [G/]GATTTCATCGACGGGTCTGT
542G6/G7-1	CTTTTAGGCCTTGACAAATC [A/G] CAATCCGGCCCCTTTTGATC
S42G6/G7-2	CTGCACTTGTGCAGACCATC[A/G]CCCTGATCGCTGTCAGGGAG
S42G5/G7-1	CTTTGCTCTGCACTTGTGCA (A/G) ACCATCGCCCTGATCGCTGT
S42G5/G7-2	TCGCTGTCAGGGAGGCGTC[T/G]TGCAGCTCGGCTCCATGAAA
S42G5/G7-3	CAGCTCGGCTCCATGAAAAA(G/)GTGCCGCTACTCTCAGTC
542G2/G7-1	GTTCTCTGCACTCCGATTGA (G/A) GGTCCGAAGCAGGGCAGCGC
S42G2/G6-1	CTCCATGAAAAAGGTGCCGC[/G]TACTCTCTCAGTCAGCTACT
942G2/G3-1A	CTGCACTCCGATTGAGGGTC [C/G] GAAGCAGGGCAGCGCGTGTG
S42G2/G3-1B	CTGCACTCCGATTGAGGGTC[C/G]GAAGCAGGGCAGCGCGTTGT
542G2/G3-1C	CTGCACTCCGATTGAGGGTC (C/G) GAAGCAGGGCAGCGCGTTTG
S42G2/G3-1D	CTGCACTCCGATTGAGGGTC [C/G] GAAGCAGGGCAGCGCGTTTT
S42G1/G7-1	CAGCTCGGCTCCATGAAAAA [G/]GTGCCGCTACTCTCTCAGTC
541G2/G5-1	GATAGCCACATAGAACCTAT[G/T]CCTCCAGCTAACAACACGAG
S41G2/G5-2	CCCTCTTGACTCCTGCCGAT [T/G] GAGAAATAATTTTTTAGTAT
S41G5/G2-1	TTAGTATACCCATGACTATC(T/)TACTAAACTCTGTCAAAGCT
S41G2/G4-1	TGACTCCTGCCGATTGAGAA (A/G)TAATTTTTTAGTATACCCAT

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S41G3/G1-1	CGCCAGATGAACCAGCTCAC [TA/AGT] ATCCTGCAAGAGATATCCCT
S41G1/G3-1	TTGCAATCAGAGAACATCGA [T/C] CAATAACTGACCTTGCTTTG
S40G5/G4-1	ACAATAAATGTATTAATATA [T/C] TTTTTATAAATTTAATTGAA
S40G4/G2-1	AAGTACATGACACCTCCGGA [G/A] ATACGGCCAAGAATCCCGCA
S40G4/G1-1	AGACGCTTGAAACGCATAAG[T/]TACATGACACCTCCGGAGAT
S40G1/G4-1	TTCCGTATGGACCAATCCAT[/C]CACGGCTTTGGAGCTTAACG
S40G1/G4-2	GGAGCTTAACGATACCAATC [GA/CG] AAATTCCCTCGCCGGGCAC
S38G5/G7-1	GAGATATATGGTACTTCTAG [G/T] AGATTCAAAGACATGGAGCA
S38G5/G7-2	ATAATGATGGAACCATGACT[G/A]CCGAAGATGAACCAACCTTG
	in the state of th
S37G6/G2-1	GGTCAATAAGATAACTACAC [C/A] AAACTCTGCGTACAGTCTCG
S37G2/G6-1	GTCACTTACTCTGCTCGTGG(A/)AAGAAGTACTTGCCCTGCAG
S37G2/G5-1	GTCACTTACTCTGCTCGTGG[A/]AAGAAGTACTTGCCCTGCAG
S37G2/G4-1	TCTTATCAATTCTGTGCATT(C/T)AATACGATCAGCCAGACAGT
537G2/G4-2	TCTTATCAATTCTGTGCATT[C/T]AATACGATCAGCCAGACAGC
S37G2/Q4-3	AATTCAGCTCAAATCATAGG[T/G] CAATAAGATAACTACACAAA
S37G2/G4-4	
S37G2/G3-1	GTCTCCCGGATCCAGAATCC[C/T]ACCGCGTTACATCGCCTTTC
S37G3/G2-1	CAGCGACAACATGCATAAAA (G/T) AGCAGGTCCAATTCAGCTCA
	CCTTTCGGCCGCTCAGGGCA[/T]AACAAAATCCCCTCTACGCC
S37G3/G2-2	GCGTCACTTACTCTGCTCGT (/G)GAAAGAAGTACTTGCCCTGC
S37G2/G3-2	TGCAGGCGGCAGCCGAAGCG[/A]ATCGAAGGCCTGACCGGTCT
S37G2/G3-3	ACATCGCCTTTCCCAAACCC[G/C]CCATCTCCCCAGGCCGCGGC
S37G1/G6-1	GGGCAATAAGATAACTACAC (A/C) AAACTCTGCGTACAGTCTCG
S37G1/G6-1B	GGTCAATAAGATAACTACAC (A/C) AAACTCTGCGTACAGTCTCG
\$37G1/G6-2	AATTCAGCTCAAATCATAGG[T/G]CAATAAGATAACTACACCAA
\$37G1/G6-3	CAATACGATCAGCCAGACAG (C/T) CACTGTGCAACATCAGCAAA
\$37G1/G6-3	CAATACGATCAGCCAGACAG [C/T] CACTGTGCAACATCAGCGAC
537G1/G6-4	CTCCTTTCGGCCGCTCAGGG[C/A]AGACGAAATCCCCTCTACGC
537G1/G6-5	CGAAATCCCCTCTACGCCCG [C/T] AGTGCTCCGCTAATCGATGG
S37G1/G5-1	AAGGGAGGGTGATCTCGGGG [T/]TGTGCCAGAGGGATGGCGTC
537G1/G4-1	CAGCCACTGTGCAACATCAG (CA/AG) AAAACATGCATGAAAGAGCA
S37G4/G6-1	GTCTCCCGGATCCAGAATCC[T/C]ACCGCGTTACATCGCCTTTC
S37G4/G5-1	ATCGATGGACAAGGGAGGGT [A/G]ATCTCGGGGTGTGCCAGAGG
S37G4/G5-2	CCATCTCCCCAAGGCCGCGG[C/G]GGAAGGCGAAAACGAATGCG
S37G4/G5-2B	CCATCTCCCCAAGGCCGCGG[C/G]GGAAGGCGAAAACGATGCGA
S37G3/G4-1	CCTTTCGGCCGCTCAGGGCA [A/G] ACGAAATCCCCTCTACGCCC
S37G3/G4-2	ATCGATGGACAAGGGAGGGT [G/A] ATCTCGGGGTGTGCCAGAGG
S36G2/G5-1	TTGTCATGACCAACAGTGTC [G/A] GTGGGGTGGTCTCGTTGGAT
536G2/G5-1B	TAGTCATGACCAACAGTGTC [G/A] GTGGGGTGGTCTCGTTGGAT
636G2/G5-1C	TTGTCATGACCAACACTCTC [G/A]GTGGGGTGGTCTCGTTGGGT
S36G2/G5-1D	TTGTCATGACCAACAGTGTC [G/A] GTGGGGTGGTCTCGTTGGGT
536G1/G6-1	TGTCACTTTTAGTGGTGTT (C/T) AGACTAAGGATGCTGACATT
S36G3/G6-1	TCACTTTTTAGTGGTGTTTA [C/G] ACTAAGGATGCTGACATTCT
S36G3/G6-2	TCCTTCGAGCATGACAAGAT (G/A) TCTTCTCTTCCTTGTGCAAC
36G3/G6-2B	TCCTTCGAGCATGACAAGAT [G/A] TCTTCTCTTCCTTGTGTCAC

35G3/G6-1	TCTGAGATACATTCATTTAA (C/A) ATGTCAGATAAAGAAAACTC
35G3/G6-1B	TCTGAGATACATTCATTTAA (C/A) ATGTCAGATAAAGAAAAGTA
35G3/G6-3	TCCTTATAATATCGTTGATA(C/)CCCTGTTTGTCAGTTTGTAG
35G3/G6-4	CACCAGTAGTCTGGTGGTAC(T/)TTTTTTTTTTSSGCATGTGGG
35G3/G6-5	TGCATGTGGGCAGGTACATA (A/T) ATTAAACATTCTAGGTATTT
35G1/G4-1	CATTGAGGATTAATACTTTT (G/T) GACATGTTTGTTAATTTTTA
35G3/G6-7	TGCCGCCTCAAAATGGTTTA(A/G)ATCTTTAAGCTGTCTTCCAC
35G3/G6-7B	TGCCGCCTCAAAATGCTTTA(A/G)ATCTTTAAGCTGTCTTCCTC
35G2/G6-1	CACCAGTAGTCTGGTGGTAC (/T)TTTTTTTTGCATGTGGGCAG
535C2/G6-2	TGCATGTGGGCAGGTACATA[G/T]ATTAAACATTCTAGGTATTT
535G2/G6-3	GGTTCGCACCATATCATGAT (C/T) GGAATGCCGCCTCAAAATGG
53502/G3-1	CCTACTTGTAGAACAATAGA (G/A) GTGTTCTCTACCTCTGAGAT
535G2/G3-2	GCTCGAACTTCTAGTTGATT (A/G) CATGATTGCTATTACTGTTG
835G4/G5-1	GGTTCGCACCATATCATGAT [C/T] GGAATGCCGCCTCAAAATGG
63.4G2./GE 3	CATGCCTCTGTTGATATTTT (G/C)GTGCACCTTTTGCTTGCAAC
S34G3/G5-1	GATTTTGTAGGTTGATGCAT (C/T) GTTTGATCTTTCTTATCTCC
534G3/G5-2	TGCTTGCAACTAAATTAATC [A/G] TGCTCTATTTGACTAAGAGT
S34G3/G5-3	ACATGTCCAGGACGCATGGT[C/]CCCAATATTGTTGTTGGAAG
S34G3/G4-1	TTGATCTTTATCTCCTT(/C)CGAATTTGTTCTGTTATA
S34G3/G4-2	TTGATCTTTCTTATCTCCTT[/C]CGAATTTGTTCTGTGTATAC
S34G3/G4-2B	TGTAGGACTTGGAGAGCTTG[A/G]TAATTTACACATGCCTCTGT
S34G2/G5-1	CATGCCTCTGTTGATATTTT[G/C]GTGCACCTTTTGCTTGCAAC
534G2/G5-2	GATTTTGTAGGTTGATGCAT (C/T) GTTTGATCTTCTTATCTCC
534G2/G5-3	GAGACATTTCCTACTCAATA [C/T] AATTATTTCATGAAATTATT
S34G2/G3-1	GAGACATTTCCTACTCAATA(C/T)AATTA
	AGTATCACAGACTAATCTGA (A/G)TATCTGGTTGCCACGAAAAC
S33G5/G6-1A	AGTATCACAGACTAATCTGA(A/G)TATCTGGTTGCCACAAAAAC
533G5/G6-1B	TCAAAGTGGTGCAATCGCAA [T/C] CCACTTGGGCTTGCCGTGGT
533G5/G6-2	
533G5/G6-3	CCACTTGGGCTTGCCGTGGT [C/]CGTATCGTACGCAGGTAGCA AGCATTTTTGTTTTTTTTT [T/C]CCTTGGCAGACAACAGACAG
533G5/G6-4	CAGTCCCGAGAATCCCAAAT[C/]CAGAAAAAGGTTTTGTTTTT
S33G5/G6-5A	
S33G5/G6-5B	CAGTCCCGAGAATCCCAAAT[C/]CAGAAAAAGGTTTTGTTTTA
\$33G4R/G6-1A	GGCAGACAACAGACAGATCA [AG/CA] CATGCTTGCATTTACTCCCA
S33G4R/G6-1B	GGCAGACAACAGACAGATCA (AG/CA) CATGCTTGCATTTACTCTCA
S33G3R/G6-1A	GTGATCACAGACTAATCTGA(A/G)TATCTGGTTGCCACGAAAAC
S33G3R/G6-1B	GTGATCACAGACTAATCTGA [A/G]TATCTGGTTGCCACAAAAAC
S33G3R/G6-2A	TCTGAATATCTGGTTGCCAC (G/A) AAAACCGGGACACAAGAGAG
S33G3R/G6-2B	TCTGAGTATCTGGTTGCCAC [G/A] AAAACCGGGACACAAGAGAG
533G3/G6-3	TCAGTCAAACTCAGTCCCGA(A/G)AATCCCAAATCAGAAAAAGG
S33G3/G5-1A	GGTTGCCACGAAAACCGGGA [C/G] ACAAGAGAGAAACTCAGAGT
S33G3/G5-1B	GGTTGCCACGAAAACCGGGA[C/G]ACAAGAGAGAAACTCAAAGT
S33G2R/G3-1A	ACGCATGCTTGCATTTACTC[C/T]CAGTCAAACTCAGTCCCGAA
S33G2R/G3-1B	ACACATGCTTGCATTTACTC[C/T]CAGTCAAACTCAGTCCCGAA
S33G1R/G2-1	TATTATTCAATTTTGAATAA (/G) GAAGGAAATTTTAGCACCTC
S32G3/G5-1	ATTAXTAXATGCATCCTCTG(C/G)TAAAAAAACCCATTTTGAAT
S32G3/G5-1	ATGAATTGAAGCTCTGAATA [C/T] AGAATCCACCATTCTTCCGA
532G3/G5-2 532G3/G5-3	GAATCCACCATTCTTCCGAA [A/G]CTGCTTCCTACAAAACTCGA

32G3/G5-4	GAAAGGATGTGTTTTGATA [G/A] CCTTCAGTCTTTCAGATGGA
	The second secon
31G3/G5-1	CAATGTCTTGTTCGTTATCA [A/G] CGAAAGTTTGAATCCCCACA
31G3/G4-1	TGTATCGGCTAGTCTGGATG [G/A] TCGCACTGGCACTCAGTGCT
	TCTATTCAGCAGTCTGAGAA (GCA/CT) AGGATGGTCGGCTTCTTCAG
529G4/G5-1	CCTTACACTATTAACAGGCC [C/T] GTGATCTACCTGAATGCCTG
329G1/G5-1	CCTTACACTATTAACAOOCCTCTTTTTTTTTTTTTTTTT
28G5/G6-1	CAAGAAGCCTCTTCAGTGTC [A/C] GTCGTAGCTTCCTCAAGACC
528G5/G6-2	AGACCTTCCTGATGTGCGGA [T/C] GCTAATCCATGGAGCAGGGA
528G5/G6-2B	AAGACCTCCTGATGTGCGGA[T/C]GCTAATCCATGGAGCAGGGA
528G5/G6-3	CTAATCCATGGAGCAGGGAG [G/A]AAGGGGCGAGGGGCAGCAAG
\$28G4/G5-1	TCGTCCCGAATACACCCGGG[G/C]GAGGGGGTGGTCGCGACTGG
528G3/G6-1	GTCGTAGCTTCCTCAAGACC[T/]TCCTGATGTGCGGACGCTAA
S28G3/G4-1	GAGTCGTCGCGAATACAGCC[A/G]GGGGAGGGGGGGGGGGGGGAC
S28G3/G4-2	AGGGGTGTCGCGACTGGA [T/G] CGCCCGAGCAGCGAGCAAGC
S28G3/G4-2 S28G3/G4-3	AAGCACATGTTTTAACCTTT (T/G) ATTCAAACTTTCCAGCCGTT
	AAGCACATGTTTTAACCTTT[T/G]ATTCAAACTTTCCAGCGTTA
S28G3/G4-3B	GAATGTTGCTGTTATATTAC(T/C)CGTAGGTGACAAAGGGTTCA
S28G2/G6-1	AGAAAAATTTACATAAAAAA [G/C] CACACTCCATGATTGTTAAA
S28G2/G4-1	AGAAAAATTTACATAAAAA [G/C]CACACTCCATGATTGTTTAA
528G2/G4-1B	CTTTTATTCAAACTTTCCAG[/C]CGTTAATTTGTTATCCGTTG
\$28G2/G3-1	TGTTGAACATGCTCTCAGGA[/CC]CCCCTATTGTGACACAGCA
528G6/G1-1	TACATCTTAACAAGCACATG[TG/TTT]TAACCTTTTATTCAAACTTT
S28G1/G3-1	TACATCTIAncandenento(13/111/11110000000000000000000000000000
S27G3/G6-1A	AGTAATGTGTGACTGTGGGC [C/G] CGTGTGACAGCTTTTACGTA
\$27G3/G6-1B	AGTAGTGTGACTGTGGGC [C/G] CGTGTGACAGCTTTTACGTA
S27G3/G6-2	TTCGCTTGGTAGCCGTAGCA [G/A] TATACTTTTACCGGCCACAG
\$27G3/G6-3	GGGCTTTGGGTTGTGAACTT[CCA/C]AAAAAAAAAAAAAATTTCCC
<u> </u>	
S26G5/G6-1	CCAAGAAAGATTAATGCTGG[/T]TAAAATATTGTTTCCAGTCT
S26G5/G6-2	AAAATCAGGACTGCGAAAAA (A/C)CCAAGAAAGATTAATGCTGG
S26G5/G6-2B	AAAATCAGGACTGCGAAAAA (A/C) CCAAGAAGATTAATGCTGGT
S26G5/G6-3	AAAGTGTGTTGTTGCCCA (G/A) ATGATTCCATTCCACACAAG
526G4/G5-1	AGGACTGCGAAAAACCAAG[/A]AAGATTAATGCTGGTAAAAT
526G4/G5-2	ATGCTGGTAAAATATTGTTT[/C]CAGTCTTTCACAAAGTGTGT
\$26G3/G4-1	CTACAAAATCAGGACTGCG[/A]AAAAACCAAGAAGATTAATG
· · · · · · · · · · · · · · · · · · ·	TTGTTTCAGTCTTTCACAAA [/GT]GTGTGTGTGCCAGATGATTC
S26G3/G4-2	TCACACACCGACCTGCCTGG[/T]TATCAGGAACCATCCTCCTG
\$26G3/G4-3	
S23G4/G5-1	GGTGAATTGGTGATGCATGC[T/G]GGGGGTGCTCGAGTTGGATG
S23G4/G5-2	TTCCAGTCGGATGAACTGGA[T/G]GTTCGTCATCCACTCGTCAC
S23G3/G6-1	GGTGAATTGGTGATGCATGC (A/T)GGGGGTGCTCGAGTTGGATG
S23G5/G3-1	TTAAGTGAAGATGCCCAAAC [C/G] GTTAAACTTTCCATGGAACT
523G5/G3-1B	ATTAATGAAGATGCCCAAAC [C/G]GTTAAACTTTCCATGGAACT
\$23G1/G6-1	TGATTCGGGTCTGTATGCGA [G/T] TGTTGTGGTGGTGAACTGGT
	CGGGTCTGTATGCGAGTGTT[G/A]TGGTGGTGAACTGGTGAATT
S23G1/G5-1	4.10

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S22G5/G6-1	CAGATTGGTGTCGTTTACTA [A/G] AATTCAGTTCTGTCCATTTG	
S22G5/G6-2	AAGTAAGCATTCTTTATATG[/T]TACTTCCCATGATAAACTTT	
S22G5/G6-3	CAAAGGGCTTACTGTACTTT[/C]CATCTTATTGGCAGGCCACC	
S21G3/G7-1	AGGCACTGACGGTGCGATGG[/T]TTTGACCTTAATTGGCATTC	
S21G3/G6-1	GTTCTCGCTGCCGTCACAAG [A/]ATTCTAGTAAGGTGTGGCTT	
S21G3/G6-2	ACTGATCTCAATTATGCATG[/A]AGAATGTTTCTCGCATCAAG	
521G2/G7-1	TTGACGGTGTTAGTTTCGAG (A/G) AGATAACACACGCTGGGAAT	
521G2/G7-2	CGATTATTAGCACGCTGGGA [G/T]TTGACCTTGAGCTCCAGGGA	
S21G2/G5-1	TGGCTTGGTTGACCTTGAGG [C/G] CCACACACTATCTAGTACGT	
S21G2/G5-2	TCCAAGTCATCTGCTAACTG[A/]ATCTCAATTATGCATGAGAA	
521G2/G5-2B	TCCAAGTCATCTGCTAACTG(A/)ATCTCAATTATGCATGAAGA	
S21G2/G3-1	AGTTTGACGGTGTTAGTTTC(G/A)AGAAGATAACACACGCTGGG	
s21G2/G3-2	AGAATGTTTCTCGCATCAAG(T/)TGCTAGAGCTGGAAAACGAA	
S20G5/G6-1	GCCGCCGAGAGCGAGGCATA (G/T) GCGCATGTGCATGTGCCTGC	
S19G5/G6-1	ACTTGGCCGGGGACGTCGAC (G/A) ATCGTCGTAGCACTACTGGT	
S19G5/G6-2	AGTACATGGCGAGCGTTCTA [G/C] CAGCTGCTTAGGTGATGTGG	
01005 (07.1)	GCTCATCACTTTCTTTCCAC(C/T)GTTTTTTTAGATGTGCACCG	
S17G5/G7-1A	GCTCATCACTTTCTTTCCAC (C/T) GTTTTTTTAGATGTGCACCT	
\$17G5/G7-1B		
S17G4/G7-1A	CTGTTAATACTTCTATTTCC [A/C] AGCTAACAACCCCTCTTGGT	
S17G4/G7-1B	TGTTTAATACTTCTATTTCC[A/C]AGCTAACAACCCCTCTTGGT	
S17G4/G7-2	CCTCTTGGTCCCAACATCCT[G/T]GAAAACTTCGGAAAAGGCTT	
S17G4/G6-1	CTATTTCCAAGCTAACAACC[C/G]CTCTTGGTCCCAACATCCTG	
817G3/G7-1	TOGGCCTAACCAGTGATTTT [A/T] GGCTGAATCTTGTCTTGTGC	
S17G3/G7-2	TCCTCTTGTGCTGTTACTGT [C/T] TTCTCTACTGCCTGTACTGA	
517G3/G7-2	TGCACCTATGTGTCTTATTA (A/T) CATGCGATTCTACTAGTTTA	
517G3/G7-3	AAGCCTCATGTGCAGATTCA (TC/AG) GAACACACAACGTCAGCCAT	
S17G3/G7-4	ACAACGTCAGCCATGAGCCC [C/A]TGACACAAAGAATCTGCACT	
S17G3/G6-1	GOTTCTAAACATAGCTCGTC [C/A]ATTCATGATTCATCTCGAGC	
<u> </u>	The second secon	
S16G6/G7-1		
S16G4/G6-1	TCAGCAAGCCTCCAAGGCTC [C/A] AATGGTCCAGTTACTTGGTT	
S16G2/G7-1A	TGCTTTCGTAAACCTAGAGT (T/G) GACCAGCTGTGATTTCGGTG	
516G2/G7-1B	TGCTTTCGTAAACCTAGAGT[T/G]GACCAGCTGTGATTTCGATG	
S16G2/G7-2A	GTTGACCAGCTGTGATTTCG [G/A]TGTATTCCACGACCACGAGT	
S16G2/G7-2B	GTGGACCAGCTGTGATTTCG (G/A) TGTATTCCACGACCACGAGT	
S16G2/G6-1	GTGTGTAGCTTCATTCGCAA (TG/AT)TTTGAACAGCCTCTGCAAGT	
S16G2/G6-2A	GTGCTTTCGTAAACCTAGAG [T/C] TGACCAGCTGTGATTTCGGT	
S16G2/G6-2B	GTGCTTTCGTAAACCTAGAG (T/C) TGACCAGCTGTGATTTCGAT	
S16G2/G6-3A	GCTGACCAGCTGTGATTTCG [G/A] TGTATTCCACGACCACGAGT	
S16G1/G7-1A	CTTAATTGTACACAGTGCTT [C/T] CGTAAACCTAGAGTTGACCA	
S16G1/G7-1B	CTTAATTGTACACAGTGCTT [C/T] CGTAAACCTAGAGTGGACCA	
S16G2/G7-2B	TGCTTCCGTAAACCTAGAGT(T/G)GACCAGCTGTGATTTCGATG	
S16G1/G6-1	TGTGTAGCTTCATTCGCAAA (G/T) TTTGAACAGCCTCTGCAAGT	
S16G1/G3-2A	GTGCTTCCGTAAACCTAGAG (T/C)TGACCAGCTGTGATTTCGAT	
S16G1/G3-2B	GTGCTTCCGTAAACCTAGAG(T/C)TGACCAGCTGTGATTTCGGT	

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S16G1/G2-1	GTGTGTAGCTTCATTCGCAA (A/T)GTTTGAACAGCCTCTGCAAG
514G5/G7-1	CTGGACCAGATCGCCCTGCC[/TC]TCAGATTCAGAGATTGACGA
S14G5/G7-2	CGAAAGAGCGAGATATATCG[A/G]TCGATCGATGAGCAAGTATA
514G5/G6-1	GCTCAGCTGCCGGAGTACGT (A/T)GGCTTGCTCTCCGGCCGGCC
S14G5/G6-1B	ATAGCTCTGCCGGAGTACGT (A/T) GGCTTGCTCTCCGGCCGGCC
S13G5/G6-1	TTTCACAACTCAACTGATTG(A/T)CTTGCTTTGATGTGGATTCT
513G2/G6-1	TTGGTAATTTCAGAGCTAGA (C/G) AACTTACTGTGGTACACGCC
S12G4/G6-1	ACCTTTGCTGTTTTTTTTT[T/G]GTATTCGAATGGAGGGAGTA
S12G2/G5-1	AAAACAGCCAAGGTGGTGGT [C/G] AAAGGAAGGTGTCAGAAGGT
S12G2/G5-2	TCTGTTCGTTCCATCTCTTT[A/G]CAGTAAATATCCGTAATTAC
S12G2/G5-3	CGTAATTACTTTGTTACTAC [TA/C]AGTAATTTTATATATATCCT
S12G2/G5-4	TATATATATCCTCATTTCAA [A/T] GAACAGTCAAAGTTAGTTTT
S12G2/G5-4B	TATATATCCTCATTTCAA(A/T)GAACAGTCAAAGTAGTTTTG
S12G2/G4-1	TATTTCTTATCCAGGATTGT[T/C]CTTTGGCCAAAGCATGGTAC
S12G2/G4-2	CGTTCCATCTCTTTACAGTA(A/G)ATATCCGTAATTACTTTGTT
512G2/G4-3	ATCCGTAATTACTTTGTTAC [TA/AC]CTAAGTAATTTTATATATAT
S12G2/G3-4	GTAATTACTTTGTTACTACT [A/]AGTAATTTTATATATATCCT
S12G1/G6-1	CTGTGTTTTTTTTGGTATT[G/C]GAATGGAGGGAGTATTATTT
512G1/G6-1B	GCTGTGTTTTTTTGGTATT [G/C] GAATGGAGGGAGTATTATTT
S12G1/G5-1	ACTTAGATGATGACCAGGTG[A/]AGAGTTTGGCACCTTTGCTG
S12G1/G5-2	AGTTTGGCACCTTTGCTGTG[T/]TTTTTTTTTGGTATTGGAATG
S12G1/G5-3	CTTTACTGATTGGGTTACAA (A/G) AGGTTATTTCTTATTCAGGC
S12G1/G5-4	AATTACTTTGTTACTACCAG[T/]TAATTTTATATATATCCTCC
<u> </u>	
S10G4/G6-1	AGCGACAGGGATGTCGAGCA [G/T]CTACGGAAGGCAATAATGAG
S10G4/G6-2	AATTTGGGAAAATCAATGCA [GAA/CAC] ATCAGTGATTAATCCACATA
S10G3/G6-1	GCATGCCGGAGTGAGGGAGG[TG/]TGTGTGTGTGTGGCTCCACA
S10G3/G6-2A	GCCCCTACGCCATTTAGCG[G/A]ATTTGGGAAAATCAATGCAG
S10G3/G6-2B	GGCCGCTACGCCATTTAGCG[G/A]ATTTGGGAAAATCAATGCAC
\$10G1/G6-1	CATCCCGGCGGCAGAACAA [C/G]GTACGAGAAGGATGGAATGC
S08G5/G6-1	GTCCCAGATCAGGTCCACGT[T/C]CGAGCTCGCTGTTCCCGCTT
S08G5/G6-2	TGGTTCTTCACCACCACCGC (C/G) CCGGGCGCGCCCAGCGCCTC
S08G4/G6-1	GCAGCCTCAGGTACACGGGG [/ A] AAGTCGGAGTGGTTCTTCAC
S08G4/G6-2	GCCGGGCGCCCAGCGCCT[/C]CGTCCCAGATCAGGTCCACG
S08G3/G6-1	GCACGTCGTTGGTGAAGAAG [AC/CA]GCGGTACGGGTGCTTGTCGA
S08G3/G5-1	AGGTACACGGGGAAGTCGGA [G/T] TGGTTCTTCACCACCACCGC
S08G3/G5-2	CGACGGCGTCCAGCACCGAC (G/)CCTCCGCCTTCACCCCGCGC
S08G3/G4-1	GTCCACGTCGAGCTCGCTGT (C/T) CCCGCTGCCCACGACGGCGT
S08G1/G4-1	GCACGTCGTTGGTGAAGAAG (A/C)AGCGGTACGGGTGCTTGTCG
S06G2/G3-1	NAACCAAACCCTGACTATTA (T/C) AGGTAGATTAGACTAGACAC
506G2/G3-2	ACGGTGAGGAGTGGCACATG (A/C)GATGGAAAGTTCCTGTAGAC
S06G2/G3-2E	ACGGTAAGGAGTGGCACATG (A/C) GATGGAAAGTTCCTGTAGAC
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S05G2/G4-1	TATGCTTGGAAAGTGGGAAA [G/]CGGAACATACGATGGAGGAC

	AAACAATAATTTTTACACAG[/T]TGCTAAGGTTTTACTGTTTT
03G5/G6-1	AAACAATAATTTTTACACAG(/T)TGCTAAGGTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT
03G2/G6-1	ATATCCATGTTGTCGCCTGC (/TG) TGTGCGCTTGCTTGCCGCTA
02G6/G7-1	CACGGCAGTTGGCAGTGTGG (A/)AAGGACTATCTCGTGGTGAT
02G6/G7-1	GTOGTGATGCCAACATGCAA (G/)TTACCCAATCCAGGCTTCCC
02G4/G7-1	TGGTGTTTGTTAGGAAAGAC (T/C)TAATTTCACGGCAGTTGGCA
02G4/G7-2	AATAATATTTGGCGGTGAAA(G/)CCTGGTGGAAAACTGTTTTC
02G4/G7-2B	AATAATATTTGGCGGTGAAA[G/]CCTGGTGGAAAACTGTTTCC
02G4/G6-1	TTGCTGCTATGTTTACTGGG[/T]TGTAGAAAAAAAAAAATAATAT
02G3/G7-1	CAACGTGCAATAATAGAACA(T/G)GTGGTGTTTGTTAGGAAAGA
502G3/G7-1 502G2/G6-1	GCTCGGTAATAATTCTGGCT [C/G] CGATGGCACCCATATTCCTC
502G2/G6-1B	GCTCGGTAATAATTCTGGCT (C/G) CGATGGCACCCATATTCCTG
502G2/G5-1 _	AAAACACGTGGTGTTTGTTA(G/A)GAAAGACCTAGTTTCTCGGC
S02G2/G5-1B	AAATCACGTGGTGTTTGTTA [G/A]GAAAGACCTAGTTTCTCGGC
	TAGTTTCTCGGCAATTGGCA [G/T] TGTGGAATGACCATCTCGTG
502G2/G5-1	TAGTTTCTCGGCAATTGGCA [G/T]TGTGGAATGACCATCTCGTC
S02G2/G5-1B	GTOTGGAATGACCATCTCGT [G/C] GTGATGCCAGCATGCTGTTA
S02G2/G5-2	GTGTGGAATGACCATCTCGT[G/C]GTGATGCCAGCATGCTACTA
S02G2/G5-2B	ACCCTGTCAGGCTTCCACAG (A/C)TATAATATTTGTTGTGGTGT
502G2/G5-3	ACTCTGTCAGGCTTCCACAG (A/C)TATAATATTTGTTGTGGTGT
S02G2/G5-3B	ACTCTGTCAGGCTTCCACAG(A/C)TATAATATTTGTTGTGTGTG
S02G2/G5-3C	ACCCTGTCAGGCTTCCACAG(A/C)TATAATATTTGTTGTGTGTG
S02G2/G5-3D	WCC1010V00C11carport(ii) of
	GAAARGACGAGAGAGATTCC [G/A] AAAGCCCCATCTCCCCGACT
S01G7/G4-1	GGAAAGACGAGAGAGATTCC (G/A) AAAGCCCCATCTCCCCGACT
S01G7/G4-1B	AACAGCGCAAGAGTCAACAC (A/G) CGCGCGCATCACGCATGCCA
S01G4/G7-2	AACAGCGCAAGAGTCAACAC (A/G) CGCGCGCATCACGCATACCA
S01C4/G7-2B	GCTAGAGCAAGAGTCAACAC [G/A] CGCGCGCATCACGCATGCCA
S01G4/G7-3	GCTAGAGCAAGAGTCAACAC [G/A] CGCGCGCATCACGCATACCA
S01G4/G7-3B	CACACGCGCGCATCACGCAT[G/A]CCACGAGCTTTCAGGTTATG
S01G4/G7-4	CACACGCGCGCATCACGCAT (A/G) CCACGAGCTAGCTTTCAGGT
S01G4/G7-4B	CACGCCGCCCATTGG (/A) AAAGACGAGAGATTCCGA
S01G3/G7-1	
S01G3/G7-2	AGCCAGGTTCTAACAGCTAG (C/A) GCAAGAGTCAACACGCGCGC

TABLE 3: PRIMERS PAIRS FOR AMPLIFICATION OF FRAGMENTS CONTAINING POLYMORPHISMS

NUM	FORWARD_PRIMER	REVERSE PRIMER
1	TCTCGGATGCTAAAAAGCAAGC	TCGTGGGTTGATCTCGAGGAC
2	TCTGACATCTCTCAACGTCCTT	GAAAGTCTTCAGGGATCTTCCG
3	GGTGAGATCTTAAACACCTTAAAC	CGTTTTTAACTTTTGGTTAGATTG
4	CCCAATCCCCAAATTTAAAGCA	AGGGTACCAAAGGGTACATCAT
5	ATAGCAACTACTGATAGTGAACCA	ACTCTCTTTGCTTTAAATTTGGG
6	TCTCCCAACGTATTCTCTTGGT	TCTGCTCCCCTGCAACATTAAC
7	TCAGAGGCAAAGCGGTAGTTG	GTTGGGAGATTACCTCGTGAGA
8	TCCACTACCAGCTTCAATGTCT	CGTAATGAACTTTGGCATCGAC
9	CGATGCCAAAGTTCATTACGAT	TCGTGGTCTCCATGGATGACT
10	TCTCCTTACTTCCTTTCTCCC	ATGAGTTTCTGCACTTTTCACT
11	TGACAATTTTGTTTTGTTTCTTG	TACAGAACTTTGTTCTGACCTGA
12	TGCTTTAAAAACCAGAGCCACC	TTTGGAAAGAAAGGAAACGATG
	GGCAAACAATCTAAGCGGTC	CGAGCAAGAAGCATGAGACA
13	CCATGGTAGGACAGGCAAAC	AGTGAGACTGCGGTCGAAA
14	TCTAAGCAGTGCAACAGCTCCT	AGAGCTTCCTCGTACGTCAGGG
15	CAACAAACTCAAAAAAACGATACCT	AGCGAGACGGCGAGAGACG
16		TGAAGTTCGAGATCCTTGCGAC
17	TGTTGATCAGGAGACAGTTACA	TGCGACTCTGTGGGACTGTA
18	GCTGCTCCTGATCTTGAAGG	TGCGACTCTGTGGGACTGTA
19		
20	GATAAGGTCCATGATGACA	GATCAGGTCCATGATGACA
21	CCATTGGTATACATCCAACTTT	TGATGACAAAAAGGTATTCCA
22	AGGTCCATGATGATGACAAAAA	AAAGAACTGGAGAAAAGTTGGA
23		
24		
25	AAGGACACTCCACTTTCTTCGT	AAAGACATGAGACCGGCTCG
25	AAGGACACGCCACCGTCTTC	AAAGACATGAGACCGGCTCG
25	GACACTCCACTTTCTTCGTGGC	agatgaaagacatgagaccggc
26	TCTGCTGCAAGAGATGGTGTGT	TGTGAGTATCCATCGAAGAGGT
26	CATCGCTGTTATCATCAGTACC	AGAGGTAGTAAGAATTAGCCTTGG
27	TCTGCTGCAAGAGATGGTGTGT	TGTGAGTATCCATCGAAGAGGT
27	GCTGGTGTGTTGGGTATCTACG	ATTAGCCTTGGGGTTAATACCG
28	•	
29	GTTCTTGGTCTTCCTTACGCCT	GGAGGACACCCATGAGCCAG
30	CTGGACTATGCGGTGGCT	GATAGGAGCGAACGTGAAGG
31	GACTAGTGCGGTGGCTGC	GATAGGAGCGAACGTGAAGG
32	GACTAGTGCGGTGGCTGC	GATAGGAGCGAACGTGAAGG
33	AAACGGTACTGCGAAAGCTG	GGCACCACATCGAAGTTTCT
	CCTGAAACGGTACTGCGAAAG	AAGTTTCTCAAGTTGTGGAATTC
34	CAACAACAAGCTGGTCAGGA	GCTTGAACGGAGCAGCAT
35	GTTTCTCGTCAAAAGCCACG	TCCAAGGCGCAAGAAGAG
36		TCCAAGGCGCAAGAAGAG
37	ACAACCTCGCGGTGGTAAC	AGCTTGTTGTTGGAAGCGTT
38	AAACCAGGGTCCTTGATGTG	
38	TAAGACCAGTAAGCGGAAAACC	CTTGTTGTTGGAAGCGTTGTAG
39		
40	CTTTGGGACCGTTGGAGTT	TTTCCACCGTCATAGCCG
41	GATGGTTGTGGTAGCGACTGC	TTAAGGCTGCTTCCAGGCTC
41	GATGGTTGTGGTAGCGACTGC	TTAAGGCTGCTTCCAGGCTC
42	CTGATATGGAGGTTGAAGGAGG	CTTATTTGTCTCCGGTAGCTCG
43	TACGGTCCCATCAGTGACAA	GGCTCATTCCAGCCATTTC
44	GCTGCTCTAGGGATGCTCAG	TGTCACTGATGGGACCGTAA
45	CTAGGGATGCTCAGCACCATC	TGGGACCGTAAGCATCAATC
46	ATATGATGAGAAAGTGCTTGTGG	ATGTTGCGGTGCACGAGATT
47	ATATGATGAGAAAGTGCTTGTGG	ATGTTGCGGTGCACGAGATT
47	ATATGATGAGAAAGTGCTTGTGG	TTGTAGCGGTGCACGAGATT
48	ATATGATGAGAAAGTGCTTGTGG	TTGTAGCGGTGCACGAGATT
49	TGGGAATGATTGGTTTGAGA	CGATATATCAAGCGTACCAGC
50	# # # # # # # # # # # # # # # # # # #	
51	GGAGAAACCCTCTCCGACTT	GCCGAAGGTCATGGAGAAAG

52	GTTATCGATCGCGTGGTCC	GGTGTCGGACAGGGATACTG
53 54	GTTATCGATCGCGTGGTCC	GGTGTCGGACAGGGATACTG
5 4 55	GIIMICUIIGGGGGGGG	
56 56		•
57		
58	AAAAACGAAACTGTTGCCTCA	GGCTACACCTTCATACCATGAAATA
59	TGGGTCTTGATGGATTGTTTC	CAGCTACCTCCATCAGCTTCT
60	TACCGTGCGAGAACAGAAGAGA	TCCTTAAGTACTGTCCGACGTG
61	AAACGTGTGACACAACGAAG	ACGGAACCCACGGCATTTAC
61	AAACGTGTGCCACAACGAAG	ACGGAACCCACGGCATTTAC
62	GATATCCAATACGGGTCCTAGC	AAATGCACCAGAAACGGTGAG
62	GATATCCAATACGGGTCCTAGC	AAATGCACCAGAAACGGTGAG
63	CTTGGGCTTTCGGTGGTAT	TGCAATAGACGAGGACGAAG
64	ATTAAAGCACAAAGGTCTGCCT	CGTTCCAGACACGACCAGTAC
65	CCATTTAACCTCCTTGATTCC	ATAGTTAGAAAGACCCCCAGCC ATAGTTAGAAAGACCCCCAGCC
65	CCATTTTAACCTCCTTGATTCC GATCGAAGACGAGAGCTTGAA	GTATGAAGGTTTTCCAAATCCG
66	ACTAAAACGAGAGCTTGAATAACG	TGAAGGTTTTCCAAATCACCGT
66 67	GGAGCAAAGCTAAAAAGATCGA	CTCCAAGCTTGATAGATGGAGC
68	GTCCAATGGAAGGTGGGTTTAG	CCCACAGTGAAGTTTGTATCCA
69	TGGAAGGTGGGTTTACCGT	CCAAAGATGGTTCCCACAGT
70	ATTGGGTCTTGATGGATTGTTT	GACAGCTACCTCCATCAGCTTC
71	TGGATTGTTTCTATCCTGCAAA	AGCTACCTCCATCAGCTTCTA
72	TCTACCACGGTCGTACTGGTC	GCTTGGTGACGTTCCAGAC
73	TGGTTCTGTCTCATTTTGTTGG	CCATCAATATCTACGCTTTCGA
74	TGGTTCTGTCTCATTTTGTTGG	CCATCAATATCTACGCTTTCGA
75	GCCAAAACACTGTTGGAGGAC	GTGTCCCAGATGTTGTCTTGAA
75	CCGAAAACACTGTTGGAGGAC	GTGTCCCAGATGTTGTCTTGAA
76	ATTACCCCCGTCTTTCGTATCT	TAAGGTATTTGCATCTGATCGG
77	AATGAGATCATAGTATCCGCCG	TAGTTGAGATGATGGCCCGAC
77	AATGAGATCATAGTATCCGCCG	TGTAGTTGAGCTGATGGCCC
78	AATGAGATCATAGTATCCGCCG	TAGTTGAGATGATGGCCCGAC
78	AATGAGATCATAGTATCCGCCG	TGTAGTTGAGCTGATGGCCC
79	TGCACCAACATTGTGAACCT	GTTGAAGGTGGCTGAGGAAG
80	ATGETTGCTCCAACATTGTGAA	AGCTCTTCGTTGAAGGTGGCT
80 80	ATGAAGTGTTGCTGCCACATAG	GTTGTTAGGTGGCTGAGGAAGA
81	TTGCTGCCACATAGTGAACC	TAGGTGGCTGAGGAAGAAGC
81	ATGGTTGCTCCAACATTGTGAA	AGCTCTTCGTTGAAGGTGGCT
81	ATGAAGTGTTGCTGCCACATAG	GTTGTTAGGTGGCTGAGGAAGA
82	TGCACCAACATTGTGAACCT	GTTGAAGGTGGCTGAGGAAG
82	ATGGTTGCTCCAACATTGTGAA	AGCTCTTCGTTGAAGGTGGCT
82	CAGGTTTGTTTCTACACCGTCA	GAGGCCACTGTACTCACAACAA
82	ATGAAGTGTTGCTGCCACATAG	GTTGTTAGGTGGCTGAGGAAGA
83		
84	TGTATCGGTGCCTCTATCCG	GTTGAAGGTGGCTGAGGAAG
85	GCCTCCTCAACTTGACCCTC	TTCACATCAGCGAGTCCTTG GCTTGGGGTTGGTTTAGGA
86	GCTGCCCTCTCCCAATTC GGAGACGGATAAACATGGAAAG	TACCATTCCCTCTTGCTTTGAT
87 87	GGAGACGGATAAACATGGAAAG	CCCCTTTGCTTTGATGACTAG
88	GGGGTACAAATCCTTTTTCGTT	TAATGGGTTCCACTTGGAGAAA
89	CTTGAATCTTCCTCTGCATTCC	GCTAACTCTTCCTCCGTTGCTA
90	ACCATGGTCGCTCCATTCAC	GCTAACTCTTCCTCCGTTGCTA
91	CCGAGTTGACTCAGCTTTCTTA	GAGTGAAGGAATGGATTCGTGT
91	CTGACGAGTTGACTCAGCTTTC	GAGTGAAGGAATGGATTCGTGT
92	ATTGCGGCTAACATCTCTGG	TACGGCTGGGTTAAGGTGA
93	TCCGAGGATCACTTCTCTCTG	CCTTGCCGAACTTCTTCTTG
93	ATCTGAAGAGGAGAACCCACTG	CCTGAGATTGGATTTGAAAAC
93	GGGTGTCGTAGTTTACGGAAGA	GGATTTGAAAACGGAGAATCTG
94	CCTCCGAGGATCACTTCTCTC	CGGAGAATCTGTAGTCGTGCT
94	ATCTGAAGAGGAGAACCCACTG	CCTGAGATTGGATTTGAAAAC
94	GGGTGTCGTAGTTTACGGAAGA	GGATTTGAAAACGGAGAATCTG
94	GATCACTTCTCTGTTCAAGAA	GGAGAATCTGTAGTCGTGCTCC

?	AGCTGGATCCCTCTGCTCGT	AAACTCAGACCGAGTCAGATCC
95	AAGCTGGATCCCTCTGCTC	AGACCGAGTCAGATCCGAGA
96	CAAGAAGATTCGGCAAGG	TGAGATTGGATTTGAAAACGG
97	AAGCTTCTTACTCCACTCGCC	GCGAGCCATTTCAGACAAGT
97	CCGACTCTCGAAGCTTCTTACT	TTTCACGACACGTGGACAAC
98	TCTACTTGTCTGAAATGGCTCG	AGAGGAAGATCTGCTCTCCGAG
98	AAACACGACAAACCGTTGTCTA	AGAGGAAGATCTGCTCTCCGAG
99	ACTACACCGACTCTCGAAGCTT	CGAGCCATTTCAGACAAGTAGA
100	GTCTCCGGTGAATCTAGGAGAG	TCATTCTTGTCACGACATTGAA
101	AAGCTTCTTACTCCACTCGCC	ATTTCACGACACGTGGACAA
102	TGCGTCATGAAGATAATTCACTA	CGGCGAGTGGAGTTAAGAAG
103	GGATGGTCATTCTCTTTGGTGT	ATCCATTAACTACAATGAAAAGGG
103	AGGAAGGTCATTCTCTTTGGTG	ATCCATTAACTACAATGAAAAGGG
104	CTCCTCCACGTAATCTCTCCTC	AAAGAATGTATCCGTGACGTCG
105	CTAGCTCTGGCAAGGAAAAGTG	TGAGCACATTAGCGCAGACAC
105	CAAGGGATAATTGCCCTAAAGA	TGAGCACATTAGCGCAGACAC
106	CTTTGAGAGGTTCTTCCCACAG	CGGTCAGATACTTTGGAAGGAA
107	TCATTTATGTATCAGGACAATGTG	TCTGTATTGAATGGATGCAACC
108	ACGCAACTGCCACCAATCAG	CCTCCATCACATCTTCAATGTC
109	TTTCCTCCTTTTATCCACAGGT	GAAAAGAGAGATGGGTACACAC
110	TATTATCGTTGAGGCTAGGGCA	GGGAGTGGGACCAAGATCCA
110	AGACTTGAGATGGCTGCGTATT	GGGAGTGGGACCAAGATCCA
111	TCCCTGGATCCATCTCAACTAT	AGAAAATGTCCTAGCCTCAACG
111	GTCCTGGACCATCTCAACTATG .	GTCTAGAAATGCCCTAGCCTCA
112	GCAATTTGTGTATGTGATTCGG	TAATCTCTCCTGTGGGAGGAAG
113	AAGACGCTAAGGTAGTTGGAGG	AGCATCGTGGAGATCCTTAGTC
114	GGCATGTACTCAACGTAGCAAG	GACACTCTCAGACGATTTATTTCG
115	TATCCCGAGACTTTCCCCAT	GCTGGGTTTCGACGGTGGATGG
116	CTCACCATCGGAGAAAGAGG	TCATCACTCTCATTAACAAGATCC
117	GAACAACATCTCTTCCCTCACC	GGACCTTCCCACAACAAAAATA
118	TAGGATGCGTGTGCTAAGGC	GTTACCCTTGACACGCATGTAC GTAGTGGACGATGGCTTGGAG
119	TCTTCAAAGCTTCTTACGAGCC	GTAGTGGACGATGGCTTGGAG
120		GAGAGTCCGGTTAAAGCCTTG
121	AGGACACAGAGGCTCGAAGTTA	AACTTCTTCTTGCCAACAGGAG
122	CCCAAAATAAATTGATGGTTTT	AGGAGGCCACACCTGTTTATTA
122	CCCAAATAAATTGATGGTTTT	AGTCTCAGCAGAGCGAGATTTT
123	CCGAAGAGTTTCCAGTGAAAAG	AGTCTCAGCAGAGCGAGATTTT
123	CGAAGGAGTTTCCAGTGAAAAG	TTATCCACACAATCGCTAGCAC
124	CATCTGAAGCAGCTCTGACTTG	ITATCCACACAATCGCTAGGAG
125		
127		
128		
129 130	ATCAGGAAGCCCATGGTACG	CCGGAGAATGGACCCAAGTACT
131	AICAGGAAGCCCAICCIACC	
132	ACAACTGCTGTATGATGGTAAGC	TCACATGTTCCACACAAAACAA
133	AGTCTGACTTCAAGGACTTCGC	GACGAGAGCAGAGGTTGCTAAC
133	AGTCTGACTTCAAGGACTTCGC	AAATCCGGCGATCTTGACAG
134	VOICIONCII	
135	GAGCCTGAAGGAGAGTTGGA	TGGCGATGTGTTTCATCTTC
136	CTTGGGTCCATTAGTGAGGG	TGATTTTCCTCCAATCATCAAA
137	GAAGATGAAACACATCGCCA	TCACCAACAGGATCTCCCAT
138	ACTCATAGGCGATCTGGAGTA	CTGGAGAGGAGATTCGATTTGC
139	TCTTCTGCACGCCTCACTTGTT	TTTTCCGGCGAATCCTTGAT
140	GTGTTTTTGAGGTGAAAG	AAAAAGGTATCTCCATTTGT
141	CCCGAGCCATTAGGACAAG	TTCTTACCGTGGCTTCAACC
142	CTGCACCCATCTCATCCTTT	CTGTCCCAGGAGCTTGATTC
143		
144		
145	TGAAGGCGTCGACTATCTTG	GAACTGTACGAACTGTGGCG
146	CGCCACAGTTCGTACAGTTC	CCAGTGCATAACAGTGGCAA
147	ACAATCTGAAGGCGTCCACT	CCAGTGCATAACAGTGGCAA
148	TCTTCACGGAAATCTTGCTG	CAGGCTTGTTTCGTTCAACA
149	CCTGCCCATGTCTTCAAAGT	GTCTTTCTTGTTCGGACGTG

		TCAAAAATAAATGTTAAAAATAAGAGGACA
150	GGATATTACATAATACTTGATATGCCATT GGATATTACATAATACTTGATATGCCA	AAGCCTATTCATATTTTCAAGTTAAT
150	GGATATTACATAATACI IGATATGCCA	CAGGGTATCCAGCCGTTG
151	GAAGTTCGAGACGCTGTCGT	GAGCCCAAGGAACGGAGCGAC
152	GTGATTATGGCCACGTCGGCC	TGAAGGGACAACCCAATCAT
153	AAAGTGCCGGTAATGTAATGGT	
154	GGTTCGCCTCATGTTATCGTTG	GGTGGAGGTGGGGTTTATCCAA
154	ACCACTGACGTAGCACCTCC	GGTGGGGTTTTACAACGAGA
155	ACCACGAAGAAGGGAGGTTC	GTCGGAGGTGCTACGTCAGT
156	CCAATTTATCGAGCCAGACA	TGGTCCAAGCAGTTCATCAC
157	ATGCAACATCGAGCATTCTG	TCACATCCAGAGCTGACACG
158	TGGTTGCTTACTGAGAGAATGC	GACACGTGAGCAGACTCCAA
159	TGCGCTGTGCTTTATTGGGAA	CAGATTCCATTCAGACCCTAGAGG
159	CAGTGCGCTGTGCTTTATTG	CCATTCAGACCCTAGAGGCA
160	GGAGTACACTGTTACAATCACAC	AGCTGTCTATCTGTGTTGCGCTT
160	GCGCAACACAGATAGACAGC	CGGGTCTGAAGAAAGTGGA
161	GGAGTACACTGTTACAATCACAC	AGCTGTCTATCTGTGTTGCGCTT
162	AGCGAGATCGATCCTGTTG	TCCTATGGCAGTGATGGATG
163	CGAGCAGATGGGTGCTAGTC	TGCTTTCTTAGTGGGTGAACG
164	TGAGCAAAGCAGAGACTCCA	ACCCATCTGCTCGAAGTAGC
165	AATACACGCTGCCCAGGCC	GATCTCGCTCGGTGGTTCC
165	GAATACACGCTGCCCAGG	GGCAGTGATGGATGACAACA
166	TACTAGCTCGGTGACCCGAT	ATGTGGCGAGCTTCTTTCAT
167	CAAGCAAGCAAGCTGTCTGT	GCGCGCAAACCTTTTATTAC
168	GAATCTGACCCACCCGTTGC	AAAACCAAGAAAAGCCAACATT
	TGCGCGCAATAAATCAGA	TGGGTCAGATCAGATCAGC
169	GATCTGAATCTGACCCACCC	TAACGATGAGCAAGTGCAGC
170	TCGACGTGGAAAGTATTGGA	ACAGCTGCCACTGAATCCAT
171	GCTCCTCGTGGTAGTGACGA	GACCAGAAACGCCTTGAGTC
172		TCACTACCACGAGGAGCGCC
173	CCGCCGAGGGAATACTCGAT	TCGTCACTACCACGAGGAGC
173	GCGAGGGAATACTCGATAGG	TGGCTGCTGGTGGGC
174	GATATGGGGTTCGGTGTCACTGA	GGCGCTGCTGCTGAT
174	GTTCGGTGTCACTGACCTGT	GCAGCTGCGTTATCCTCAAG
175	AAATCCGCGGAAACAACAT	CAMCING A CTCTCA A CA A A TA A CATTC
176	TCATCACTTGATGATATGAGTATATTCAA	CATGGTCAACTGTGAAGAAATAACATTC
176 177	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG
176 177 178	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG
176 177	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG
176 177 178	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT
176 177 178 179	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG
176 177 178 179 179	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTCCGGC
176 177 178 179 179 180	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTCCGGC ACGTGAAGATGGTCAGACAGA
176 177 178 179 179 180	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCAGTACAGCTA CCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTCCAGGC ACGTGAAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCGA
176 177 178 179 179 180 180	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTCCGGC ACGTGAAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCGA ATAATGAACCAGCAGGACGCC
176 177 178 179 179 180 180 181	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCAGTACAGCTA CCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTCCGGC ACGTGAAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCGA ATAATGAACCAGCAGGACGC ACGCGAGCAGTTGTGGGTTACC
176 177 178 179 179 180 180 181 182	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTCCGGC ACGTGAAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCGA ATAATGAACCAGCAGGACGC ACGCGAGCAGTTGTGGGTTACC ATAATGAACCAGCAGGACGC
176 177 178 179 179 180 180 181 182 182 183	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTCCGGC ACGTGAAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCGA ATAATGAACCAGCAGGACGC ACGCGAGCAGTTGTGGGTTACC
176 177 178 179 179 180 180 181 182 182 183 183	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTCTGGC ACCGTGAAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCGA ATAATGAACCAGCAGGACGC ACGCGAGCAGTTGTGGGTTACC ATAATGAACCAGCAGGACGC TCGAAGGACGACTTGAATCTCTC AAACAGACCCGTCGATGAAA
176 177 178 179 179 180 180 181 182 182 183 183 184	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTCCGGC ACCTGAAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCA ATAATGAACCAGCAGGACGC ACGCGAGCAGTTGTGGGTTACC ATAATGAACCAGCAGGACGC TCGAAGGACGACTTGAATCTCTC AAACAGACCCGTCGATGAAA GAGCTCTTCGGCGTAGTAGG
176 177 178 179 179 180 181 182 182 183 183 184 184	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTCCGGC ACCTGAAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCA ATAATGAACCAGCAGGACGC ACGCGAGCAGTTGTGGGTTACC ATAATGAACCAGCAGGACGC TCGAAGGACGACTTGAATCTCTC AAACAGACCCGTCGATGAAA GAGCTCTTCGGCGTAGTAGG
176 177 178 179 179 180 180 181 182 182 183 183 184 184 184	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTCTGGC ACCGTGAAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCGA ATAATGAACCAGCAGGACGC ACGCGAGCAGTTGTGGGTTACC ATAATGAACCAGCAGGACGC TCGAAGGACGACTTGAATCTCTC AAACAGACCCGTCGATGAAA
176 177 178 179 180 180 181 182 182 183 183 183 184 184 185	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTTCCGGC ACCGGAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCGA ATAATGAACCAGCAGGACGC ACGCGAGCAGTTGTGTGGGTTACC ATAATGAACCAGCAGACGC TCGAAGGACGACTTGAATCTCTC AAACAGACCCGTCGATGAAA GAGCTCTTCGGCGTAGTAGG GTAGCGGCACCTTTTTCATGG
176 177 178 179 180 180 181 182 182 183 183 184 184 185 186 186	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCAGTA CCTTGATTGCAGTTA GCGTCCTGCTGGTTCATTATCTG GCGTCCTGCTGGTTCATTAT CTGGTGGTGGTGCTTCTCT CTGCGTGCATGTGAACATTCATG ATCGACGGGTCTGTTTTCCT TGCGTGCATGTGAACATTCATG GATTTCATCGACGGGTCTGT CCGACTACCCGGAGCTGAACC GACTACCCGGAGCTGAACC GACTACCCGGAGCTGAACC TGATCGCTGTCAGGAGGGC CTGATCGCTGTCAGGGAGGC CTGATCGCTGTCAGGGAGG CGTTCTCTGCACTCCGATTGA	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTCCAGGC ACGTGAAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCGA ATAATGAACCAGCAGGACGC ACGCGAGCAGTTGTGGGTTACC ATAATGAACCAGCAGACGC TCGAAGGACGACTTGAATCTCTC AAACAGACCCGTCGATGAAA GAGCTCTTCGGCGTAGTAGG GTAGCGGCACCTTTTTCATGG CTTTTTCATGGAGCCGAGC
176 177 178 179 180 180 181 182 183 183 184 184 184 186 186	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTCCAGC ACGTGAAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCGA ATAATGAACCAGCAGGACGC ACGCGAGCAGTTGTGGGTTACC ATAATGAACCAGCAGACCC TCGAAGGACGACTTGAATCTCTC AAACAGACCCGTCGATGAAA GAGCTCTTCGGCGTAGTAGG GTAGCGGCACCTTTTTCATGG CTTTTTCATGGAGCCGAGC ACACACGCGCTGCCCTGC
176 177 178 179 180 180 181 182 182 183 183 184 184 185 186 186 186	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCAGTACAGCTA CCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTCCGGC ACGTGAAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCGA ATAATGAACCAGCAGGACGC ACGCGAGCAGTTGTGGGTTACC ATAATGAACCAGCAGGACGC TCGAAGGACGACTTGAATCTCTC AAACAGACCCGTCGATGAAA GAGCTCTTCGGCGTAGTAGG GTAGCGGCACCTTTTTCATGG CTTTTTCATGGAGCCGAGC ACACACGCGCTGCCCTGC GCTGCCCTGCC
176 177 178 179 179 180 180 181 182 182 183 183 184 185 186 186 186 187 188	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCAGTACAGCTA CCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTCCAGC ACGTGAAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCGA ATAATGAACCAGCAGGACGC ACGCGAGCAGTTGTGGGTTACC ATAATGAACCAGCAGACCC TCGAAGGACGACTTGAATCTCTC AAACAGACCCGTCGATGAAA GAGCTCTTCGGCGTAGTAGG GTAGCGGCACCTTTTTCATGG CTTTTTCATGGAGCCGAGC ACACACGCGCTGCCCTGC
176 177 178 179 179 180 180 181 182 183 183 184 185 186 186 187 188 189 190	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTGTAGCTG ATCCAGTTGATCTTCCGGC ACGTGAAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCGA ATAATGAACCAGCAGGACGC ACGCAGCAGTTGTGGGTTACC ATAATGAACCAGCAGGACGC TCGAAGGACGCAGCAGCACC TCGAAGGACGCTTGAATCTCTC AAACAGACCGTCGATGAAA GAGCTCTTCGGCGTAGTAGG CTTTTTCATGGGCTAGTAGG CTTTTTCATGGACCCTGC GCTGCCTTGCCT
176 177 178 179 179 180 180 181 182 182 183 184 184 185 186 186 186 187 188 189 190 190	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCAGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTCCGGC ACGTGAAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCGA ATAATGAACCAGCAGGACGC ACGCAGCAGTTGTGGGTTACC ATAATGAACCAGCAGGACGC TCGAAGGACGCAGCAGCACC TCGAAGGACGCTTGAATCTCTC AAACAGACCGTCGATGAAA GAGCTCTTCGGCGTAGTAGG CTTTTTCATGGGCTAGTAGG CTTTTTCATGGCCTTCC GCCGCCTGCCTGC TCGGCAGGAGCACC TCGGCAGGAGCACC TCGGCAGGAGCACCTTCATGGC GCTGCCTTGCCT
176 177 178 179 180 180 181 182 182 183 183 184 184 185 186 186 186 187 188 189 190 190	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCAGTACAGCTA CCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTCCGGC ACGTGAAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCGA ATAATGAACCAGCAGGACGC ACGCGAGCAGTTGTGGGTTACC ATAATGAACCAGCAGGACGC TCGAAGGACGACTTGAATCTCTC AAACAGACCCGTCGATGAAA GAGCTCTTCGGCGTAGTAGA GAGCTCTTCGGCGTAGTAGG GTAGCGGCACCTTTTTCATGG CTTTTTCATGGAGCCGAGC ACACACGCGCTGCCCTGC GCTGCCTTGCCT
176 177 178 179 180 180 181 182 182 183 184 184 185 186 186 187 188 189 190 190 190 191	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCAGTACAGA TGACCTTGATTGCAGTA CCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTGATCTTCCGGC ACGTGAAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCGA ATAATGAACCAGCAGGACGC ACGCGAGCAGTTGTGGGTTACC ATAATGAACCAGCAGGACGC TCGAAGGACGACTTGAATCTCTC AAACAGACCCGTCGATGAAA GAGCTCTTCGGCGTAGTAGG GTAGCGGCACCTTTTTCATGG CTTTTTCATGGAGCCGAGC ACACACGCGCTCCCTGC GCTGCCCTGCC
176 177 178 179 180 180 181 182 182 183 183 184 184 185 186 187 186 187 188 189 190 190 191 192 193	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCAGTACAGA TGACCTTGATTGCAGTA CCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCACTGTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTCCGGC ACGTGAAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCGA ATAATGAACCAGCAGGACGC ACGCGAGCAGTTGTGGGTTACC ATAATGAACCAGCAGGACGC TCGAAGGACGACTTGAATCTCTC AAACAGACCCGTCGATGAAA GAGCTCTTCGGCGTAGTAGG GTTACTCGGCGTAGTAGG CTTTTTCATGGAGCCGAGC ACACACGCGTCCCTGC GCTGCCCTGCC
176 177 178 179 180 180 181 182 182 183 183 184 184 185 186 187 188 189 190 190 190 191 192 193	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCAGTACAGA TGACCTTGATTGCAGTACAGCTA CCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCCACCG GGCCTTCATCTAGTCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTCCGGC ACGTGAAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCGA ATAATGAACCAGCAGGACGC ACGCGAGCAGTTGTGGGTTACC ATAATGAACCAGCAGGACGC TCGAAGGACGCTCGATGAATCTCTC AAACAGACCCGTCGATGAAA GAGCTCTTCGGCGTAGTAGG GTAGCGGCACCTTTTCATGG CTTTTCATGGACCCGACC ACACACGCGCTGCCCTGC GCTGCCCTGCC
176 177 178 179 180 180 181 182 182 183 183 184 184 185 186 186 187 188 189 190 190 191 192 193 194 195	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCAGTACAGA TGACCTTGATTGCAGTA CCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTCCGGC ACCAGTGATCTTCCGGC ACGTGAAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCGA ATAATGAACCAGCAGGACGC ACGCGAGCAGTTGTGGGTTACC ATAATGAACCAGCAGGACGC TCGAAGGACGCTTGAATCTCTC AAACAGACCCGTCGATGAAA GAGCTCTTCGGCGTAGTAGG GTAGCGGCACCTTTTTCATGG CTTTTTCATGGAGCCGAGC ACACACGCGCTGCCCTGC GCTGCCCTGCC
176 177 178 179 180 180 181 182 183 183 184 185 186 186 186 187 188 189 190 190 191 192 193 194 195 196	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCAGTACAGA TGACCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCAGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTCCGGC ACGTGAAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCGA ATAATGAACCAGCAGGACGC ACGCGAGCAGTTGTGGGTTACC ATAATGAACCAGCAGGACGC TCGAAGGACGCTTGAATCTCTC AAACAGACCCGTCGATGAAA GAGCTCTTCGGCGTAGTAGG GTAGCGGCACCTTTTTCATGG CTTTTTCATGGAGCCAGGC ACCACGCGCTGCCTGC GCTGCCTTGCTTCGGATAT GCGCAGGAGTCAAGAGGGATAT GGCAGGAGTCAAGAGGGATA GCGCTCGGCATATTTAACTCGTGT TCTGGATGGGTCCCGGC CTCGAAGCTGATCCAAGGTTG CCCAGTGCTCCAAGGTTGAA AAGGATAGGCTGGTCCTCGAAC ACGCGCTGCCCTGC CCCAGTGCTCCAAGGTTCCCCCC ACGCGGTGCTCCATGTCTTTGAA AAGGATAGGCTGGTTCCCTCC ACGCGGTGGGATTCTTGGAA AAGGATAGGCTGGTTCCCTCC ACGCGGTGGGATTCTTGGAA
176 177 178 179 180 180 181 182 182 183 183 184 184 185 186 186 187 188 189 190 190 191 192 193 194 195	TCATCACTTGATGATATGAGTATATTCAA GCTGCGTCAATCATCACTTC CCACAGGCGTCAAGTACAGA TGACCTTGATTGCAGTACAGA TGACCTTGATTGCAGTA CCTTGATTGCATTGC	CATGGTCAACTGTGAAGAAATAACATTC TCTGTACTTGACGCCTGTGG TGTCAGACTCCAAGGCGAAG TGATCTTCCGGCCTTCCACG GGCCTTCATCCAGTTGATCT CTTCCACGGGCTTCTTGTAGCTG ATCCAGTTGATCTTCCGGC ACCAGTGATCTTCCGGC ACGTGAAGATGGTCAGACAGA GATAATGAACCAGCAGGACGCGA ATAATGAACCAGCAGGACGC ACGCGAGCAGTTGTGGGTTACC ATAATGAACCAGCAGGACGC TCGAAGGACGCTTGAATCTCTC AAACAGACCCGTCGATGAAA GAGCTCTTCGGCGTAGTAGG GTAGCGGCACCTTTTTCATGG CTTTTTCATGGAGCCGAGC ACACACGCGCTGCCCTGC GCTGCCCTGCC

199	GTGCATTCAATACGATCAGC	TGAATTGGACCTGCTCTTTT
	AAAGAGCAGGTCCAATTCAG	GACTGTACGCAGAGTTTTGTGTAG
200		GAAAGGCGATGTAACGCGG
201	GATCGAAGGCCTGACCGGT	
201	GAAGCGATCGAAGGCCTGA	TGGGAAAGGCGATGTAACG
202	ACCATGTTAATGTCACTTTTTAGTGG	GATATATTCCCTACGAGGAATGTC
		ATATTTGTCCCTAAACCTCGGGATAT
202	CATTGTGAGGAaCACCATGTTAATG	
203	GGATTAATACTTTTGGACATGTTTG	ATATGGTGCGAACAACTCAACAGTA
203	TTTGACATGTTTCTTAATTTTTATAG	GAACCCTCAACAGTAATAGCAATCAT
	- -	TATAAGGATAAATGTCCTACTTTTCT
204	TCTACCTCTGAGATACATTCATTTAA	
205	GAGGGTTCGCACCATATCATG	CCATTTTGAGGCGGCATTCC
205	TATTACTGTTGAGGGTTCGCAC	AACCATTTTGAGGCGGCATT
	TGGCCATTGTAGATATAATATGAT	TCGAAGGAGATAAGAAAGATCAAA
206		- -
207	GGGGATATTATGTAGGACTTGGAG	CACCAAAATATCAACAGAGGCATG
208	CCTTGGCAGACAACAGACAGATCA	TTTGACTGGGAGTAAATGCAAGC
	TGGCAGACAACAGACAGATCA	GGGATTTTCGGGACTGAGTT
208		
209	TTGTTTTGTTTTTCCTTGGCAGAC	TTCGGACTGGGAGTAAATGCA
210	ACTCAAAGTGGTGCAATCGCAA	CGATACGGACCACGGCAAGC
210	ACCGGGAGACAAGAGAGAAACT	GATACGGACCACGGCAAGC
	**	GGGATTTTCGGGACTGAGTTTGAC
211	AGACAGATCAGTAAATCACGCATG	
211	AGACAGATCAGTAAATCACGCATG	TTGGGATTTTgGGGACTGAG
212	TTTGAATGAATTGAAGCTCTGAA	TTTCAATTGTTGCATCTCGAGTT
	TTTTGAATGAATTGAAGCTCTG	TCTCGAGTTTTGTAGGAAGCAG
212		
213	ATCCTCTGCTAAAAAAACCCATT	TCGAGTTTTGTAGGAAGCAGTTT
214	ACCGGACTCCTTTCCAATGTC	TCAGTATGTGGGGATTCAAAC
215	AATGGCAACTTGTATCGGCTAG	CATGTTACCTATTAGAGCACTGAG
		GTACTGGCTGAAGAAGCCGA
216	GAGTGCAGAACCTGATGGGT	
217	TGCAGTCAGGGTTTTTTTTTCAAC	AGCTCAGGCATTCAGGTAGATC
218	CCTTGTGGAATGTTGCTGTTAT	ACGACGGACACTGAAGAGGCT
219	TCTTCAGTGTCCGTCGTAGC	CCCTTTCTCCCTGCTCCAT
		GCTTGCTCGCTGCTCGGGC
220	CCGGGCGAGGGGGTGGTC	
220	CAGGGGAGGGGGTGGTCGC	GTGTGCTTGCTCGCTGCTC
221	TCCGAGCCCACAAATCTGTAC	CAAATTAACGCTGGAAAGTTTGAA
	ATATTACCCGTAGGTGACAAA	CAGGAAGGTCTTGAGGAAGCT
222		
222	CGTAGGTGACAAAGGGTTCAA	AGGAAGGTCTTGAGGAAGCTA
223	AGCTTGTCTTCGCTTGGTAGC	TTCACAACCCAAAGCCCTG
223	AGCTTGTCTTCGCTTGGTAGC	CCTGTGGCCGGTAAAAGTATA
224	GCGAAAAAACCAAGAAAGATTA	ATCTGGGCAACAACACACACTTT
225	CTGCGAAAAACCAAGAAGATTA	ATCATCTGGCACACACACTTTG
226	TGAGAACAATGAGACACACGAATT	GTACAGAATTGTGGAGTTCCATG
		GTACAGAATTGTGGAGTTCCATG
226	CGAATTAAGTGAAGATGCCCAAA	GIACAGAMI IGIGGAGII CCAIG
227	GCCTGATTCGGGTCTGTATGC	TGCATCACCAATTCACCAGTT
228	TGACTCGGCTCAATTCCAG	ATGAAAGTGACGAGTGGATG
229	CGGTAGCTATGCACAGATTG	TGCATTCAAATGGACAGAACTG
230	CTAGTAAGGTGTGGCTTGGTTGAC	CGAAACTAACACCGTCAAACTCTA
230	GTGGCTTGGTTGACCTTGAGG	CGAAACTAACACCGTCAAACTCTA
231	TCCGAACGCCGCCGAGAGC	GGCAGGCAGGCACAT
	TCCGAACGCCGCCGAGAGC	GGCAGGCAGGCACAT
231		
232	GTGAGGCGAAGTACATGGCG	AGCCACATCACCTAAGCAGC
233	GCGCGTCGGTTCTAAACATA	AACCAGCTGTGCTCGAGATGA
	ACTTCTATTTCCAAGCTAACAACC	TTTCCAGGATGTTGGGACC
234		
235	TAAATTAAAGCCTCATGTGCA	TTCTTTGTGTCAGGGGCTCAT
236	GGTGTGTGCTCCAAGTGTGT	AAGCACTTGCAGAGGCTGTT
237	TCTAACTCAGCAAGCCTCCAAG	CTACACACTTGGAGCACACACC
		ACTCGTGGTCGTGGAATACA
238	GTTGACCAGCTGTGATTTCG	WCICGIGGICGIGGWINCA
239	GCCTCTGCAAGTGCTTAATTGTA	GTCGTGGTCGTGGAATACATCG
239	TTGAACAGCCTCTGCAAGTG	CATCGAAATCACAGCTGGTC
240		
	3 C3 C3 CTCCTTCCCTT3 3 5 CC	CAMPEA AAMEACACCOCCOC
241	ACACAGTGCTTCCGTAAACC	CATCGAAATCACAGCTGGTC
242	CACAGTGCTTTCGTAAACCTAGA	TGGTCGTGGAATACATCGAA
243	ATGAACGAAAGAGCGAGATA	GGCCGTATACTTGCTCATCG
244	AATAGCTCAGCTGCCGGAGT	AGCAGCAGCAGTGCATGG
		TCGAATGGCGTGTACCACAGTA
245	GGTTGGTAATTTCAGAGCTAGA	TCGAATGGCGTGTACCACAGTA
246	CGATGGGTTTTCACAACTCA	GTAACCTAGAATCCACATCAAAGC
247	CCATCTCTTTACAGTAAATATCCG	GACTGTTCTTTGAAATGAGGAT

248	CCATCTCTTTACAGTAAATATCCG	GACTGTTCTTTGAAATGAGGAT
249	ATGCCTTTACTGATTGGGTTACA	GTTTTTGGAAGGAGTAATAGATT
250	CAGAAGGTCTGTTCGTTCCA	TGACTGTTCTTTGAAATGAGGA
251	GCTACGCCATTTAGCGAATTT	GTGGGAGGATAAAAAAGATATTGCT
251	AGCGAATTTGGGAAAATCAA	GAAGACAATGGTGGGAGGAT
252	TATAATGTTTAGCGACAGGGATG	TAGCTCATTATTGCCTTCCGTAG
252	TAGCGACAGGGATGTCGAG	AGCTCATTATTGCCTTCCGT
253	GCACGCACGTCGTTGGTGA	TCGACAAGCACCCGTACCGC
253	GCACGTCGTTGGTGAAGAAG	GGTGTACCCCGTCGACAA
254	GCACGTCGTTGGTGAAGAAG	GTCGACAAGCACCCGTACC
255	TCAGGTACACGGGGAAGTCGG	GCGCGGTGGTGAAGAA
255	CTCAGGTACACGGGGAAGTC	CGGTGGTGGTGAAGAACC
256	CCCAGCGCCTCGTCCCAGATC	GTGGGCAGCGGGACAGCGAG

WHAT IS CLAIMED IS:

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1. A nucleic acid segment comprising at least 10 contiguous nucleotides including a polymorphic site from a sequence shown in Table 1 or Table 2, or a plant cognate variant thereof, or the complement of the segment.

2. The nucleic acid segment of claim 1, wherein the segment is less than 100 bases.

3. The segment of claim 1 that is less than 50 bases.

4. The segment of claim 1 that is less than 20 bases.

5. The nucleic acid segment of claim 1, wherein the segment contains at least 10 contiguous nucleotides including a polymorphic site from a sequence shown in Table 1.

- 20 6. The nucleic acid segment of claim 5, wherein the sequence is a Brassica sequence.
 - 7. The nucleic acid segment of claim 5, wherein the sequence is a corn sequence.
 - 8. The nucleic acid segment of claim 1 that is DNA.
 - 9. The nucleic acid segment of claim 1 that is RNA.
- 30 10. The segment of claim 1, wherein the polymorphic site is diallelic.
 - 11. An allele-specific oligonucleotide that hybridizes to a sequence shown in Table 1 or 2, or its complement.
 - 12. The allele-specific oligonucleotide of claim 11 that is a probe.

13. The allele-specific oligonucleotide of claim 11, wherein the a central position of the probe aligns with the polymorphic site in the sequence.

- 5 14. The allele-specific oligonucleotide of claim 11 that is a primer.
- 15. The allele-specific oligonucleotide of claim 11, wherein the 3' end of the primer aligns with the polymorphicsite of the segment.

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- 16. A method of analyzing a nucleic acid, comprising: obtaining the nucleic acid from a plant, and determining a base occupying any one of the polymorphic sites shown in Table 1 or 2.
- 17. The method of claim 16, wherein the determining comprises determining a set of bases occupying a set of the polymorphic sites shown in Table 1.

18. The method of claim 17, wherein the nucleic acid is obtained from a plurality of plants, and a base occupying one of the polymorphic positions is determined in each of the subjects, and the method further comprises testing each plant for the presence of a phenotype, and correlating the presence of the phenotype with the base.

- 19. Use of a polymorphism shown in Table 1 or 2, a nucleic acid segment of claim 1, or an allele-specific oligonucleotide of claim 11 to determine common or disparate ancestry between two or more plants.
- 20. Use of a polymorphism shown in Table 1 or 2, a nucleic acid segment of claim 1, or an allele-specific oligonucleotide of claim 11 in plant breeding.
- 21. Use according to claim 20, wherein the polymorphism, nucleic acid segment or allele-specific

oligonucleotide is used to monitor a genetic contribution of an ancestral plant to a progeny plant.

- 22. Use of a polymorphism shown in Table 1 or 2, a nucleic acid segment of claim 1, or an allele-specific oligonucleotide of claim 11 to trace progeny of a proprietary plant.
- 23. Use of a polymorphism shown in Table 1 or 2, a nucleic acid segment of claim 1, or an allele-specific oligonucleotide of claim 11 in certification of a hybrid plant.
- 24. Use of a polymorphism shown in Table 1 or 2, a nucleic acid segment of claim 1, or an allele-specific oligonucleotide of claim 11 to identify a progeny of a backcross of a selected plant with an ancestral plant having a desired genetic contribution from the ancestral plant.

FIGURE 1 Brassica 18A2 - allele G

Brassica 18A2 - allele C

...TCAAAACTAATTTTTTTTTTTGATTGGTAATAAACAGGT... ccattatttg accattattt Baccattatt aaccattat ctaaccatta Actaaccatt Bactaaccat Gaactaacca agaactaacc aaGaactaac aagaactaa paaaGaacta gaaagaact ttgattata<mark>H</mark>agaaaaGaac tgattata attata tata gattata ttata ata reverse strand forward strand ... TCAAAACTAATATTTCTTTGTTGGTAATAAACAGGT... cattatttg accattattt accattatt aaccattat ctaaccatta ctaaccatt aactaaccat cactaacca **actaacc** pacaactaac haaCaactaa aaaCaacta tgattatafagaaaacaact ttgattata<mark>H</mark>agaaaaCaac gattata attata ttata tata

= perfect match = "homo" mismatch (e.g.: A-A; C-C)

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A. CLASSIFICATION OF SUBJECT MATTER			
IPC(6) :C07H 21/00, 21/02, 21/04; C12N 15/09; C12Q 1/68 US CL :435/6, 172.3; 536/23.1, 24.3			
According (According to International Patent Classification (IPC) or to both national classification and IPC		
	documentation searched (classification system followe	d by classification symbols)	
U.S. :	435/6, 172.3; 536/23.1, 24.3		
Documental	tion searched other than minimum documentation to the	e extent that such documents are included	in the fields searched
Electronic d	data base consulted during the international search (n	ame of data base and, where practicable	e, search terms used)
Databases Search Te	s:APS, Medline, Biosis, Genbank erms:Brassica; polymorph?; restriction; fragment?; le	ength; rflp; lemieux?/au; landry?/au; sa	polsky?/au
C. DOC	UMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where a	ppropriate, of the relevant passages	Relevant to claim No.
Y	CROSS et al. Purification of CpG islabinding column. Nature Genetics. May 244, see entire document, especially see	rch 1994, Vol. 6, pages 236-	1-24
Y	OHNUMA et al. Archaebacterial Ether-linked Lipid Biosynthetic Gene: Expression Cloning, Sequencing, and Characterization of Geranylgeranyl-Diphosphate Synthase. Journal of Biochemistry. 20 May 1994, Vol. 269, No. 20, pages 14792-14797, see entire document, especially sequences.		1-24
Y	AHOUSE et al. Mouse MHC Class I-Like Fc Receptor Encoded Outside the MHC. Journal of Immunology. 01 December 1993. Vol. 151, No. 11, pages 6076-6088. See entire document, especially sequences.		1-24
X Furth	ner documents are listed in the continuation of Box C	See patent family annex.	
Special categories of cited documents: T			
	"A" document defining the general state of the art which is not considered the principle or theory underlying the invito be of particular relevance		invention
	rlier document published on or after the international filing date	"X" document of particular relevance; the	
cite	cument which may throw doubts on priority claim(s) or which is ed to establish the publication date of another citation or other	"Y" document of particular relevance; the	e claimed invention cannot be
O do	ocial reason (as specified) cument referring to an oral disclosure, use, exhibition or other sams	considered to involve an inventive combined with one or more other suc being obvious to a person skilled in	step when the document is h documents, such combination
"P" doe	cument published prior to the international filing date but later than e priority date claimed	"&" document member of the same paten	
	actual completion of the international search	Date of mailing of the international se	arch report
19 MARC	CH 1998	2 9 APR 1998	
Name and mailing address of the ISA/US Commissioner of Patents and Trademarks Box PCT BRIAN R. STANTON		wB	
	n, D.C. 20231 - Io. (703) 305-3230	Telephone No. (703) 308-0196	fa

International application No. PCT/US97/21782

C (Continua	tion). DOCUMENTS CONSIDERED TO BE RELEVANT	
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Υ"	WILSON et al. 2.2 Mb of contiguous nucleotide sequence from chromosome III of C. elegans. Nature. 03 March 1994, Vol. 368, pages 32-38, see entire document, especially sequences.	1-24
Y	SAIKI et al. Analysis of enzymatically amplified β-globin and HLA-DQα DNA with allele-specific oligonucleotide probes. Nature. 13 November 1986, Vol. 324, pages 163-166, see entire document.	1-24
Y	PURDY et al. Cloning, nucleotide sequences and characterization of a gene encoding superoxide dismutase from Campylobacter jejuni and Campylobacter coli. Microbiology. 1994, Vol. 140, pages 1203-1208, see entire document, especially sequences.	1-24
Y	GUSELLA, J.F. DNA polymorphism and human disease. Annual Review of Biochemistry. 1986, Vol. 55, pages 831-854, see entire document.	1-24
Y	LOK et al. The nucleotide sequence of the 5' end of Papaya Mosaic Virus RNA: Site of in vitro Assembly initiation. Virology. 1986, Vol. 153, pages 289-296, see entire document, especially sequences.	1-24
Y	WHARTON et al. Nucleotide sequence from the neurogenic locus Notch implies a gene product that shares homology with proteins containing EGF-like repeats. Cell. December 1985, Vol. 43, pages 567-581, see entire document, especially sequences.	1-24
Y	BOARDMAN et al. Regulation of expression of a Xenopus borealis embryonic/larval alpha3 skeletal-actin gene. European Journal of Biochemistry. 1992, Vol. 208, pages 241-249, see entire document, especially sequences.	1-24

International application No. PCT/US97/21782

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
Please See Extra Sheet.
·
1. As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: 1-24 (in part)
Remark on Protest
No protest accompanied the payment of additional search fees.

International application No. PCT/US97/21782

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

This application contains claims directed to more than one species of the generic invention. These species are deemed to lack Unity of Invention because they are not so linked as to form a single inventive concept under PCT Rule 13.1. The species are as follows:

Each of the 1282 sequences listed in Tables 1-3 are a separate species because there is no requisite structural relationship between the species. Therefore, the species cited above do not relate to a single inventive concept under PCT Rule 13.1 because under PCT Rule 13.2, the species lack the same or corresponding special technical feature. There is no structural relationship between the separately recited sequences.

Applicant is entitled to have ten (10) specified sequences searched and no more than four (4) additional sequences search for each additional fee. Since no fee has been paid, the first ten (10) sequences listed in Table 1 have been searched.